# FORM PTO-1390 (REV 10-2000) US DEPARTMENT OF COMMERCE PATENT AND TRADEMARK OFF'CE TRANSMITTAL LETTER TO THE UNITED STATES

P2534-3

	DESIGNATED/ELECT	U.S APPLICATION NO. (If known, see 37 CFR 1 5)							
	CONCERNING A FILI	09/869566							
INTERNATIONAL APPLICATION NO. PCT/US99/30720		INTERNATIONAL FILING DATE 22 December 1999 (22.11.99)	PRIORITY DATE CLAIMED 23 December 1998 (23.12.98)						
TITLE	F INVENTION IL-1 RELATED	POLYPEPTIDES							
APPLICANT(S) FOR DO/EO/US GODDARD Audrey and PAN James									
Applicant	herewith submits to the United State	s Designated/Elected Office (DO/EO/US) the follow	owing items and other information:						
1.	This is a FIRST submission of item	s concerning a filing under 35 U.S.C. 371.							
2.	This is a SECOND or SUBSEQUE	NT submission of items concerning a filing under	35 U S C. 371.						
3.	This is an express request to promptly begin national examination procedures (35 U.S.C 371(f)).								
4.	The US has been elected by the expi	ration of 19 months from the priority date (PCT	Article 31)						
5.	A copy of the International App	lication as filed (35 U.S.C. 371(c)(2))							
	<ul> <li>a.   is attached hereto (requ</li> </ul>	ired only if not communicated by the Interna	ntional Bureau).						
	C-71	by the International Bureau.							
		— · · · · · · · · · · · · · · · · · · ·							
6. H		of the International Application as filed (35							
7. K	Amendments to the claims of the International Application under PCT Article 19 (35 U.S.C. 371(c)(3))								
	a.  are attached hereto (required only if not communicated by the International Bureau).								
		ed by the International Bureau.							
	_	wever, the time limit for making such amend	lments has NOT expired.						
	d. have not been made and will not be made.								
8. 📙	An English language translation of the amendments to the claims under PCT Article 19 (35 U.S.C. 371(c)(3)).								
9.	An oath or declaration of the inv	An oath or declaration of the inventor(s) (35 U.S.C. 371(c)(4)).							
10.	An English language translation of the annexes to the International Preliminary Examination Report under PCT Article 36 (35 U.S.C. 371(c)(5)).								
Items 1	1 to 16 below concern document	t(s) or information included:							
11.	An Information Disclosure Statement under 37 CFR 1.97 and 1.98.								
12.	An assignment document for recording. A separate cover sheet in compliance with 37 CFR 3.28 and 3.31 is included.								
13. 🔲	A FIRST preliminary amendment.								
	A SECOND or SUBSEQUENT preliminary amendment.								
14.	A substitute specification.								
15.	A change of power of attorney and/or address letter.								
16. 🔽	Other items or information:								
		evival Of An International Application For ally Under 37 CFR 1.137(b); fee of \$1240 sit Account No. 07-0630);							
	Enclosed is courtesy copy of International Preliminary Examination Report;								

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17. The fo	llowing fees are submitte	vd.		CAI	CULATIONS		ONLY		
BASIC NATIONAL FEE (37 CFR 1.492 (a) (1) - (5) ):									
Neither inter	national preliminary exan	nination fee (37 CFR 1.482)		l					
nor internation	nor international search fee (37 CFR 1.445(a)(2)) paid to USPTO and International Search Report not prepared by the EPO or JPO								
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Surcharge of \$136 months from the	0.00 for furnishing the or earliest claimed priority	th or declaration later than date (37 CFR 1.492(e)).	20 🗾 30	\$	130.00				
CLAIMS	NUMBER FILED	NUMBER EXTRA	RATE						
Total claims	30 - 20 -		X \$18.00	\$	180.00				
Independent claims	4 -3=		X \$80.00	\$	80.00				
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Fee for recording the enclosed assignment (37 CFR 1.21(h)). The assignment must be accompanied by an appropriate cover sheet (37 CFR 3.28, 3.31). \$40.00 per property +					40.00				
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NOTE: Where an appropriate time limit under 37 CFR 1.494 or 1.495 has not been met, a petition to revive (37 CFR 1.137(a) or (b)) must be filed and granted to restore the application to pending status.									
SEND ALL CORRESPONDENCE TO.									
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Patent Docket No. P2534-3

### IN THE UNITED STATES PATENT AND TRADEMARK OFFICE

In re Application of
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Glory L. Fabuena

### CERTIFICATE RE: SEQUENCE LISTING RESPONSE UNDER 37 CFR § 1.821(f) and (g)

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I hereby state that the Sequence Listing submitted herewith is submitted in paper copy and a computer-readable diskette and that the information recorded in computer readable form is identical to the written sequence listing. I further state that this submission includes no new matter.

Respectfully submitted,

GENENTECH, INC.

Date: February 19, 2002

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### IL-1 RELATED POLYPEPTIDES

### FIELD OF THE INVENTION

The present invention relates generally to the identification and isolation of novel DNAs having homology to interleukin-1 (IL-1) or interleukin-1 receptor antagonist (IL-1Ra) polypeptides, and to the recombinant production of novel polypeptides, designated herein as interleukin-1-like polypeptides ("IL-1lp").

### BACKGROUND OF THE INVENTION

Interleukin-1 refers to two proteins (IL- $1\alpha$  and IL- $1\beta$ ) which play a key role early in the inflammatory response (for a review, see Dinarello, <u>Blood</u>, <u>87</u>: 2095-2147 (1996) and references therein). both proteins are made as intracellular precursor proteins which are cleaved upon secretion to yield mature carboxy-terminal 17 kDa fragments which are biologically active. In the case of IL- $1\beta$ , this cleavage involves an intracellular cysteine protease, known as ICE, which is required to release the active fragment from the inactive precursor. The precursor of IL- $1\alpha$  is active.

These two proteins act by binding to cell surface receptors found on almost all cell types and triggering a range of responses either alone or in concert with other secreted factors. These range from effects on proliferation (e.g. fibroblasts, T cells) apoptosis (e.g. A375 melanoma cells), cytokine induction (e.g. of TNF, IL-1, IL-8), receptor activation (e.g. E-selectin), eicosanoid production (e.g. PGE2) and the secretion of degradative enzymes (e.g. collagenase). To achieve these effects, IL-1 activates transcription factors such as NF-KB and AP-1. Several of the activities of IL-1 action on target cells are believed to be mediated through activation of kinase cascades that have also been associated with cellular stresses, such as the stress activated MAP kinase JNK/SAPK and p38.

A third member of the IL-1 family was subsequently discovered which acts as a natural antagonist of IL-1α and IL-1β by binding to the IL-1 receptor but not transducing an intracellular signal or a biological response. The protein is called IL-1Ra (for IL-1 receptor antagonist) or IRAP (for IL-1 receptor antagonist protein). At least three alternatively spliced forms of IL-1Ra exist: one encodes a secreted protein, also known as secretory IL-1Ra ("sIL-1Ra") (described in Eisenberg et al., Nature, 343: 341-346 (1990)), and the other two encode intracellular proteins. IL-1α, IL-1β and IL-1Ra exhibit approximately 25-30% sequence identity with each other and share a similar three dimensional structure consisting of twelve β-strands folded into a β-barrel, with an internal thrice repeated structural motif.

There are three known IL-1 receptor subunits. The active receptor complex consists of the type I receptor and IL-1 accessory protein (IL-1RAcP). The type I receptor is responsible for binding of the IL-1 $\alpha$ , IL-1 $\beta$  and IL-1Ra ligands, and is able to do so in the absence of the IL-1RAcP. However, signal transduction requires the interaction of IL-1 $\alpha$  or IL-1 $\beta$  with the IL-1RAcP. IL-1Ra does not interact with the IL-1RAcP and hence cannot induce signal transduction. A third receptor subunit, the type II receptor, binds IL-1 $\alpha$  and IL-1 $\beta$  but cannot transduce signal due its lack of an intracellular domain. Instead, the type II receptor either acts as a decoy in its membrane bound form or as an IL-1 antagonist in a processed, secreted form, and hence inhibits IL-1 activity. The type II receptor weakly binds to IL-1Ra.

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WO 00/39297 PCT/US99/30720

Many studies using IL-1Ra, soluble IL-1R derived from the extracellular domain of the type I IL-1 receptor, antibodies to IL-1α or IL-1β, and transgenic knockout mice for these genes have shown that IL-1 plays a role in a number of pathophysiologies (for a review, see Dinarello, Blood, 87: 2095-2147 (1996)). For example, IL-1Ra has been shown to be effective in animal models of septic shock, rheumatoid arthritis, graft-versus-host disease (GVHD), stroke, cardiac ischemia, psoriasis, inflammatory bowel disease, and asthma. In addition, IL-1Ra has demonstrated efficacy in clinical trials for rheumatoid arthritis and GVHD, and is also in clinical trials for inflammatory bowel disease, asthma and psoriasis.

More recently, interleukin-18 (IL-18) was placed in the IL-1 family (for a review, see Dinarello et al, <u>J. Leukocyte Biol.</u>, <u>63</u>: 658-664 (1998)). IL-18 shares the  $\beta$ -pleated, barrel-like form of IL-1 $\alpha$  and IL-1 $\beta$ . In addition, IL-18 is the natural ligand for the IL-1 receptor family member formerly known as IL-1R-related protein (IL-1Rrp) (now known as the IL-18 receptor (IL-18R)). IL-18 has been shown to initiate the inflammatory cytokine cascade in a mixed population of peripheral blood mononuclear cells (PBMCs) by triggering the constitutive IL-18 receptors on lymphocytes and NK cells, inducing TNF production in the activated cells. TNF, in turn, stimulates IL-1 and IL-8 production in CD14+ cells. Because of its ability to induce TNF, IL-1, and both C-C and C-X-C chemokines, and because IL-18 induces Fas ligand as well as nuclear translocation of nuclear factor xB (NF-xB), IL-18 ranks with other pro-inflammatory cytokines as a likely contributor to systemic and local inflammation.

### SUMMARY OF THE INVENTION

A family of cDNA clones (DNA85066, DNA96786, DNA94618, DNA102043, DNA114876, DNA102044, DNA92929, DNA96787, and DNA92505) has been identified, having homology to interleukin-1, that encode novel polypeptides. The novel polypeptides and variants thereof are collectively designated in the present application as "interleukin-1-like polypeptides" or "IL-llp", as further defined herein. Accordingly, one aspect of the invention is an isolated IL-llp polypeptide.

In another embodiment, the invention provides an isolated nucleic acid molecule encoding an IL-1lp polypeptide.

In another embodiment, the invention provides a method for producing an IL-11p comprising culturing a host cell comprising a heterologous nucleic acid sequence encoding an IL-11p polypeptide, under conditions wherein the IL-11p polypeptide is expressed, and recovering the IL-11p polypeptide from the host cell.

In another embodiment, the invention provides an anti-IL-1lp antibody.

In another embodiment, the invention provides chimeric molecules comprising an IL-11p polypeptide fused to a heterologous polypeptide or amino acid sequence. An example of such a chimeric molecule comprises an IL-11p polypeptide fused to an epitope tag sequence or a Fc region of an immunoglobulin.

In another embodiment, the invention provides an antibody which specifically binds to an IL-1lp polypeptide. Optionally, the antibody is a monoclonal antibody.

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In yet another embodiment, the invention concerns agonists and antagonists of a native IL-11p polypeptide. In a particular embodiment, the agonist or antagonist is an anti-IL-11p antibody.

In a further embodiment, the invention concerns a method of identifying agonists or antagonists of a native IL-1lp polypeptide, by contacting the native IL-1lp polypeptide with a candidate molecule and monitoring a biological activity mediated by said polypeptide.

In a still further embodiment, the invention concerns a composition comprising an IL-11p polypeptide, or an agonist or antagonist as hereinabove defined, in combination with a pharmaceutically acceptable carrier.

## BRIEF DESCRIPTION OF THE DRAWINGS

Figure 1 shows a nucleotide sequence (SEQ ID NO:1) and derived amino acid sequences (SEQ ID NO:2-3) related to a native sequence hIL-1Ra1. The nucleotide sequence (SEQ ID NO:1) contains an intron believed to extend from nucleotide positions 181 to 432, with a splice donor site at nucleotide positions 181 to 186 and splice acceptor site at nucleotide positions 430 to 432. The arnino acid sequences (SEQ ID NOS:2 and 3) are derived from the exonic sequences that are believed to make up the processed (intron-free) coding sequence.

Figure 2 shows the nucleotide sequence (SEQ ID NO:4) and derived amino acid sequence (SEQ ID NO:5) of a native sequence hIL-IRal polypeptide fused at its N-terminus to a heterologous signal peptide (amino acid positions 1-15), flag peptide affinity handle (amino acid positions 16-23) and peptide linker (amino acid positions 24-36).

Figure 3 shows the nucleotide sequence (SEQ ID NO:6) and derived amino acid sequence (SEQ ID NO:7) of a native sequence hlL-1Ra1 polypeptide. The nucleotide sequence (SEQ ID NO:6) and derived amino acid sequence (SEQ ID NO:7) are believed to represent the processed (intron-free) form and intact hlL-1Ra1 polypeptide, respectively, of the nucleotide sequence (SEQ ID NO:1) and amino acid sequences (SEQ ID NO:2-3) of Figure 1. The start and stop codons in the coding sequence are located at nucleotide positions 103-105 and 682-684, respectively. The putative signal sequence extends from amino acid positions 1 to 14. A putative cAMP- and cGMP-dependent protein kinase phosphorylation site is located at amino acid positions 33-36. Putative N-myristoylation sites are located at amino acid positions 50-55 and 87-92.

Figure 4 shows a nucleotide sequence (SEQ ID NO:8) of which nucleotides 145-629 correspond to EST Al014548.

Figure 5 shows the nucleotide sequence (SEQ ID NO:9) and derived amino acid sequence (SEQ ID NO:10) of a native sequence hiL-IRa2 polypeptide. The start and stop codons in the coding sequence are located at nucleotide positions 96-98 and 498-500, respectively. The putative signal sequence extends from amino acid positions 1-26.

Figure 6 shows the nucleotide sequence (SEQ ID NO:11) of EST 1433156.

Figure 7 shows the nucleotide sequence (SEQ ID NO:12) and derived amino acid sequence (SEQ ID NO:13) of a native sequence hIL-IRa3 polypeptide. The start and stop codons in the coding sequence are located at nucleotide positions 1-3 and 466-468, respectively. The putative signal sequence extends from amino acid positions 1-33. Putative N-myristoylation sites are located at amino acid positions 29-34, 30-35, 60-65, 63-68, 73-78, 91-96 and 106-111. An interleukin-1-like sequence is located at amino acid positions 111-131.

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WO 00/39297 PCT/US99/30720

Figure 8 shows the nucleotide sequence (SEQ ID NO:14) of EST 5120028.

Figure 9 shows the nucleotide sequence (SEQ ID NO:15) and derived amino acid sequence (SEQ ID NO:16) of a native sequence mIL-1Ra3 polypeptide. The start and stop codons in the coding sequence are located at nucleotide positions 145-147 and 610-612, respectively. The putative signal sequence extends from amino acid positions 1-33. Putative N-myristoylation sites are located at amino acid positions 29-34, 60-65, 63-68, 91-96 and 106-111. An interleukin-1-like sequence is located at amino acid positions 111-131.

Figure 10 shows the nucleotide sequence (SEO ID NO:17) of EST W08205.

Figure 11 is an autoradiograph of Northern blots depicting expression of hIL-1Ra3 mRNA in placental tissue and expression of mIL-1Ra3 mRNA in day-17 mouse embryo tissue.

Figure 12 is an amino acid sequence alignment of native sequence hlL-1Ra1L (SEQ ID NO:19), hlL-1Ra1V (SEQ ID NO:25), hlL-1Ra1S (SEQ ID NO:21), hlL-1Ra2 (SEQ ID NO:10), hlL-1Ra3 (SEQ ID NO:16) polypeptides with secretory hlL-1Ra (also referred to as "slL-1Ra" and "hlL-1Ra") (SEQ ID NO:26), hlL-1Raβ (SEQ ID NO:27) and TANGO-77 (SEO ID NO:28).

Figure 13A is a Western blot depicting the interleukin-18 receptor (IL-18R) binding activity of hlL-1Ra1. In the top panel (depicting a protein band at approximately 22 kD), a conditioned medium containing FLAGhlL-1Ra1 and FLAGIL-1R-ECD-Fc (shown in the left lane) and a conditioned medium containing FLAGhlL-1Ra1 and FLAGIL-18R-ECD-Fc (shown in the right lane) were each immunoprecipitated with protein G-sepharose, and the resulting precipitates were resolved by gel electrophoresis and Western blotting with anti-FLAG monoclonal antibody. In the middle and bottom panels (depicting protein bands at approximately 22 kD and 85 kD), a second aliquot from the FLAGhlL-1Ra1 and FLAGIL-1R-ECD-Fc conditioned medium used in the top panel (shown in the left lane) and a second aliquot from the FLAGhlL-1Ra1 and FLAGIL-18R-ECD-Fc conditioned medium used in the top panel (shown in the right lane) were each immunoprecipitated with anti-FLAG monoclonal antibody, and the resulting precipitates were resolved by gel electrophoresis and Western blotting with anti-FLAG monoclonal antibody.

Figure 13B is a Western blot depicting the IL-1R binding activity of hIL-1Ra3. In the top panel (depicting a protein band at approximately 20 kD), a conditioned medium containing hIL-1Ra3-FLAG and FLAGDR6-Fc (shown in the left lane), a conditioned medium containing hIL-1Ra3-FLAG and FLAGIL-1R-ECD-Fc (shown in the middle lane), and conditioned medium containing hIL-1Ra3-FLAG and FLAGIL-18R-ECD-Fc (shown in the right lane) were each immunoprecipitated with protein G sepharose, and the resulting precipitates were resolved by gel electrophoresis and Western blotting with anti-FLAG monoclonal antibody. In the middle and bottom panels (depicting protein bands at approximately 20 kD and 85 kD), a second aliquot from the hIL-1Ra3-FLAG and FLAGIL-1R-ECD-Fc conditioned medium used in the top panel (shown in the left lane), a second aliquot from the hIL-1Ra3-FLAG and FLAGIL-1R-ECD-Fc conditioned medium used in the top panel (shown in the right lane) were each immunoprecipitated with anti-FLAG monoclonal antibody, and the

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WO 00/39297 PCT/US99/30720

resulting precipitates were resolved by gel electrophoresis and Western blotting with anti-FLAG monoclonal antibody.

Figure 14 is a Western blot depicting the interleukin-1 receptor (IL-1R) binding activity of mIL-1Ra3. In the top panel (depicting a protein band at approximately 21 kD) and the bottom panel (depicting protein bands at approximately 85 kD) the FLAGIL-1R-ECD-Fc in conditioned medium (shown in the left lane) and the FLAGIL-18R-ECD-Fc in conditioned medium (shown in the right lane) were immobilized with protein G-agarose, the resulting solid phase was contacted with conditioned medium containing FLAGmIL-1Ra3, and the resulting bound complexes were resolved by gel electrophoresis and Western blotting with anti-FLAG monoclonal antibody.

Figure 15 shows the nucleotide sequence (SEQ ID NO:18) and derived amino acid sequence (SEQ ID NO:19) of a native sequence hIL-1RalL polypeptide. The start and stop codons in the coding sequence are located at nucleotide positions 4-6 and 625-627, respectively. The putative signal sequence extends from amino acid positions 1 to 34. A putative cAMP- and CGMP-dependent protein kinase phosphorylation site is located at amino acid positions 47-50. Putative N-myristoylation sites are located at amino acid positions 64-69 and 101-106.

Figure 16 shows the nucleotide sequence (SEQ ID NO:20) and derived amino acid sequence (SEQ ID NO:21) of a native sequence hIL-1Ra1S polypeptide. The start and stop codons in the coding sequence are located at nucleotide positions 4-6 and 505-507, respectively. A putative signal sequence extends from amino acid positions 1 to 46. A putative N-myristoylation site is located at amino acid positions 61-66.

Figure 17 shows the single stranded nucleotide sequence (SEQ ID NO:23) of EST AI343258 (lower strand) along with its complementary nucleotide sequence (SEQ ID NO:22) (upper strand).

Figure 18 is an amino acid sequence alignment of native sequence hIL-1Ra1 (SEQ ID NO:3), hIL-1Ra1L (SEQ ID NO:19), hIL-1Ra1V (SEQ ID NO:25) and hIL-1Ra1S (SEQ ID NO:21) polypeptides.

Figure 19 shows the nucleotide sequence (SEQ ID NO:24) and derived amino acid sequence (SEQ ID NO:25) of a native sequence hIL-1Ra1V polypeptide. The start and stop codons in the coding sequence are located at nucleotide positions 73-75 and 727-729, respectively. An alternate start codon is located at nucleotide positions 106-108. A putative signal sequence extends from amino acid positions 1 to 48.

### DETAILED DESCRIPTION OF THE PREFERRED EMBODIMENTS

#### I. Definitions:

The terms "interleukin-1-like polypeptide", "interleukin-1-like protein", "IL-1lp", "IL-1lp polypeptide", and "IL-1lp protein" encompass any native sequence IL-1lp, and further encompass IL-1lp variants (which are further defined herein). The IL-1lp may be isolated from a variety of sources, such as from human tissue types or from another source, or prepared by recombinant and/or synthetic methods.

A "native sequence IL-11p" comprises a polypeptide having the same amino acid sequence as a native sequence hIL-1Ra1, hIL-1Ra1L, hIL-1Ra1V, hIL-1Ra1S, hIL-1Ra2, hIL-1Ra3, or mIL-1Ra3, (which are further defined herein). Such native sequence IL-11p can be isolated from

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nature or can be produced by recombinant and/or synthetic means. The term "native sequence IL-11p" specifically encompasses naturally-occurring truncated or secreted forms (e.g., a processed, mature sequence) and naturally-occurring allelic variants of the IL-11p.

The terms "naturally-occurring amino acid sequence" and "native amino acid sequence" mean any amino acid sequence found in a polypeptide existing in nature, i.e. present in a naturally-occurring polypeptide.

The terms "non-naturally-occurring amino acid sequence" and "non-native amino acid sequence" mean any amino acid sequence not found in a polypeptide existing in nature, i.e. not present in a naturally-occurring polypeptide.

"IL-1lp variant" is defined as any polypeptide that comprises a variant of hIL-1Ra1, hIL-1Ra1L, hIL-1Ra1V, hIL-1Ra1S, hIL-1Ra2, hIL-1Ra3, or mIL-1Ra3 (which are further defined herein).

Human interleukin-1 receptor antagonist analog 1 ("hIL-1Ra1"), hIL-1Ra1 polypeptide, and hIL-1Ra1 protein are defined as any native sequence hIL-1Ra1 or variant hIL-1Ra1.

A "native sequence hIL-1Ra1" means a polypeptide comprising a naturally-occurring amino acid sequence selected from the group consisting of: (1) the amino acid sequence of amino acid residues from at or about 37 to at or about 63 of Figure 2 (SEQ ID NO:5); (2) the amino acid sequence of amino acid residues from at or about 37 to at or about 203 of Figure 2 (SEQ ID NO:5); (3) the amino acid sequence of amino acid residues from at or about 15 to about 53 of Figure 3 (SEQ ID NO:7); (4) the amino acid sequence of amino acid residues from at or about 15 to at or about 193 of Figure 3 (SEQ ID NO:7); and (5) the amino acid sequence of any naturally-occurring truncated or secreted form or any naturally-occurring allelic variant of a polypeptide comprising the amino acid sequence of (1) or (2) or (3) or (4). In one embodiment of the invention, the native sequence hIL-1Ra1 comprises amino acids from at or about 37 to at or about 203 of Figure 2 (SEQ ID NO:5) or amino acids from at or about 15 to at or about 193 of Figure 3 (SEQ ID NO:7).

"hIL-1Ra1 variant" is defined as any hIL-1Ra1 N-terminal variant or hIL-1Ra1 full sequence variant (which are further defined herein).

"hlL-1Ra1 N-terminal variant" means any hlL-1Ra1 other than a native sequence hlL-1Ra1, which variant is an active hlL-1Ra1, as defined below, having at least about 80% amino acid sequence identity with an amino acid sequence selected from the group consisting of: (1) the amino acid sequence of amino acid residues from at or about 37 to at or about 63 of Figure 2 (SEQ ID NO:5); and (2) the amino acid sequence of amino acid residues from at or about 15 to at or about 53 of Figure 3 (SEQ ID NO:7). Such hlL-1Ra1 N-terminal variants include, for instance, hlL-1Ra1 polypeptides wherein one or more amino acid residues are added, or deleted, internally or at the N- or C-terminus, in the sequence of amino acid residues from at or about 37 to at or about 63 of Figure 2 (SEQ ID NO:5) or in the sequence of amino acid residues from at or about 15 to at or about 53 of Figure 3 (SEQ ID NO:7). Ordinarily, an hlL-1Ra1 N-terminal variant will have at least about 80% amino acid sequence identity, or at least about 85% amino acid sequence identity, or at least about 95% amino acid sequence selected from the group

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consisting of: (1) the amino acid sequence of amino acid residues from at or about 37 to at or about 63 of Figure 2 (SEQ ID NO:5); and (2) the amino acid sequence of amino acid residues from at or about 15 to at or about 53 of Figure 3 (SEQ ID NO:7).

"hIL-1Ra1 full sequence variant" means any hIL-1Ra1 other than a native sequence hIL-1Ra1, which variant retains at least one biologic activity of a native sequence hIL-1Ra1, such as the ability to bind IL-18R, and which variant has at least about 80% amino acid sequence identity, or at least about 85% amino acid sequence identity, or at least about 95% amino acid sequence identity with an amino acid sequence identity, or at least about 95% amino acid sequence identity with an amino acid sequence selected from the group consisting of: (1) the amino acid sequence of amino acid residues from at or about 203 of Figure 2 (SEQ ID NO:5); and (2) the amino acid sequence of amino acid residues from at or about 193 of Figure 3 (SEQ ID NO:7). Such hIL-1Ra1 full sequence variants include, for instance, hIL-1Ra1 polypeptides wherein one or more amino acid residues are added, or deleted, internally or at the N- or C-terminus, in the sequence of amino acid residues from at or about 37 to at or about 203 of Figure 2 (SEQ ID NO:5) or in the sequence of amino acid residues from at or about 15 to at or about 193 of Figure 3 (SEQ ID NO:7).

Human interleukin-1 receptor antagonist analog 1 long ("hIL-1Ra1L"), hIL-1Ra1L polypeptide, and hIL-1Ra1L protein are defined as any native sequence hIL-1Ra1L or hIL-1Ra1L variant (which are further defined herein).

A "native sequence hIL-1Ra1L" means a polypeptide comprising a naturally-occurring amino acid sequence selected from the group consisting of: (1) the amino acid sequence of amino acid residues from at or about 26 to at or about 44 of Figure 15 (SEQ ID NO:19); (2) the amino acid sequence of amino acid residues from at or about 26 to at or about 207 of Figure 15 (SEQ ID NO:19); and (3) the amino acid sequence of any naturally-occurring truncated or secreted form or any naturally-occurring allelic variant of a polypeptide comprising the amino acid sequence of (1) or (2). In one embodiment of the invention, the native sequence hIL-1Ra1L comprises amino acids from at or about 26 to at or about 207 of Figure 15 (SEQ ID NO:19).

"hIL-1Ra1L variant" is defined as any hIL-1Ra1L N-terminal variant or hIL-1Ra1L full sequence variant or hIL-1Ra1L fusion variant (which are further defined herein).

"hIL-1Ra1L N-terminal variant" means any hIL-1Ra1L other than a native sequence hIL-1Ra1L, which variant is an active hIL-1Ra1L, as defined below, having at least about 80% amino acid sequence identity with the amino acid sequence of amino acid residues from at or about 26 to at or about 44 of Figure 15 (SEQ ID NO:19). Such hIL-1Ra1L N-terminal variants include, for instance, hIL-1Ra1L polypeptides wherein one or more amino acid residues are added, or deleted, internally or at the N- or C-terminus, in the sequence of amino acid residues from at or about 26 to at or about 44 of Figure 15 (SEQ ID NO:19). Ordinarily, an hIL-1Ra1L N-terminal variant will have at least about 80% amino acid sequence identity, or at least about 90% amino acid sequence identity, or at least about 90% amino acid sequence of amino acid residues from at or about 26 to at or about 44 of Figure 15 (SEQ ID NO:19).

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"hIL-1Ra1L full sequence variant" means any hIL-1Ra1L other than a native sequence hIL-1Ra1L, which variant retains at least one biologic activity of a native sequence hIL-1Ra1L, such as the ability to bind IL-18R, and which variant has at least about 80% amino acid sequence identity, or at least about 85% amino acid sequence identity, or at least about 95% amino acid sequence identity with the amino acid sequence of amino acid residues from at or about 26 to at or about 207 of Figure 15 (SEQ ID NO:19). Such hIL-1Ra1L full sequence variants include, for instance, hIL-1Ra1L polypeptides wherein one or more amino acid residues are added, or deleted, internally or at the N- or C-terminus, in the sequence of amino acid residues from at or about 26 to at or about 207 of Figure 15 (SEQ ID NO:19).

"hIL-1Ra1L fusion variant" means a chimeric hIL-1Ra1L consisting of a native sequence hIL-1Ra1L fused at its N- or C-terminus to a heterologous amino acid or amino acid sequence. In one embodiment, the hIL-1Ra1L fusion variant polypeptide consists of a native sequence of hIL-1Ra1L fused at its N-terminus or C-terminus to a heterologous amino acid or amino acid sequence, wherein the heterologous amino acid or amino acid sequence, i.e. the resulting chimeric sequence is non-naturally occurring. In another embodiment, the hIL-1Ra1L fusion variant consists of the amino acid sequence of amino acids from at or about 26 to at or about 207, inclusive of Figure 15 (SEQ ID NO:19), or the amino acid sequence of amino acid residues from at or about 1 to at or about 207, inclusive of Figure 15 (SEQ ID NO:19), fused at its N-terminus or C-terminus to a heterologous amino acid or amino acid sequence to form a non-naturally occurring fusion protein. Such hIL-1Ra1L fusion variants include, for instance, hIL-1Ra1L polypeptides wherein a heterologous secretion leader sequence is fused to the N-terminus of the sequence of amino acid residues from at or about 26 to at or about 207 of Figure 15 (SEQ ID NO:19), or amino acid residues from at or about 1 to at or about 207 of Figure 15 (SEQ ID NO:19), or amino acid residues from at or about 1 to at or about 207 of Figure 15 (SEQ ID NO:19).

Human interleukin-1 receptor antagonist analog 1 long allelic variant ("hlL-1Ra1V"), hlL-1Ra1V polypeptide, and hlL-1Ra1V protein are defined as any native sequence hlL-1Ra1V or hlL-1Ra1V variant (which are further defined herein).

A "native sequence hIL-1Ra1V" means a polypeptide comprising a naturally-occurring amino acid sequence selected from the group consisting of: (1) the amino acid sequence of amino acid residues from at or about 46 to at or about 55 of Figure 19 (SEQ ID NO:25); (2) the amino acid sequence of amino acid residues from at or about 218 of Figure 19 (SEQ ID NO:25); (3) the amino acid sequence of amino acid residues from at or about 218 of Figure 19 (SEQ ID NO:25); and (5) the amino acid sequence of any naturally-occurring truncated or secreted form or any naturally-occurring allelic variant of a polypeptide comprising the amino acid sequence of (1) or (2) or (3) or (4). In one embodiment of the invention, the native sequence hIL-1Ra1V comprises amino acids from at or about 46 to at or about 218 of Figure 19 (SEQ ID NO:25), or amino acids from at or about 37 to at or about 218 of Figure 19 (SEQ ID NO:25), or amino acids from at or about 218 of Figure 19

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(SEQ ID NO:25), or amino acids from at or about 1 to at or about 218 of Figure 19 (SEQ ID NO:25).

"hlL-1Ra1V variant" is defined as any hlL-1Ra1V N-terminal variant or hlL-1Ra1V full sequence variant or hlL-1Ra1V fusion variant (which are further defined herein).

"hIL-1Ra1V N-terminal variant" is defined as any hIL-1Ra1V other than a native sequence hIL-1Ra1V, which variant is an active hIL-1Ra1V, as defined below, having at least about 80% amino acid sequence identity with the amino acid sequence of amino acid residues from at or about 46 to at or about 89 of Figure 19 (SEQ ID NO:25). Such hIL-1Ra1V N-terminal variants include, for instance, hIL-1Ra1V polypeptides wherein one or more amino acid residues are added, internally or at the N- or C-terminus, in the sequence of amino acid residues from at or about 46 to at or about 89 of Figure 19 (SEQ ID NO:25). Ordinarily, an hIL-1Ra1V N-terminal variant will have at least about 80% amino acid sequence identity, or at least about 95% amino acid sequence identity, or at least about 95% amino acid sequence identity with the sequence of amino acid residues from at or about 46 to at or about 89 of Figure 19 (SEQ ID NO:25).

"hIL-1Ra1V full sequence variant" means any hIL-1Ra1V other than a native sequence hIL-1Ra1V, which variant retains at least one biologic activity of a native sequence hIL-1Ra1V, such as the ability to bind IL-18R, and which variant has at least about 80% amino acid sequence identity, or at least about 85% amino acid sequence identity, or at least about 95% amino acid sequence identity with the sequence of amino acid residues from at or about 46 to at or about 218 of Figure 19 (SEQ ID NO:25).

"hIL-1Ra1V fusion variant" means a chimeric hIL-1Ra1V consisting of a native sequence hIL-1Ra1V fused at its N- or C-terminus to a heterologous amino acid or amino acid sequence. In one embodiment, the hIL-1Ra1V fusion variant polypeptide consists of a native sequence of hIL-1Ra1V fused at its N-terminus or C-terminus to a heterologous amino acid or amino acid sequence, wherein the heterologous amino acid or amino acid sequence is heterologous to the native sequence, i.e. the resulting chimeric sequence is non-naturally occurring. In another embodiment, the hIL-1Ra1V fusion variant consists of the amino acid sequence of amino acids from at or about 46 to at or about 218 of Figure 19 (SEO ID NO:25), or the amino acid sequence of amino acids from at or about 37 to at or about 218 of Figure 19 (SEO ID NO:25), or the amino acid sequence of amino acids from at or about 12 to at or about 218 of Figure 19 (SEQ ID NO:25), or the amino acid sequence of amino acids from at or about 1 to at or about 218 of Figure 19 (SEQ ID NO:25), fused at its N-terminus or C-terminus to a heterologous amino acid sequence to form a non-naturally occurring fusion protein. Such hIL-1Ra1V fusion variants include, for instance, hIL-1Ra1V polypeptides wherein a heterologous secretion leader sequence is fused to the N-terminus of the sequence of amino acid residues from at or about 46 to at or about 218 of Figure 19 (SEQ ID NO:25), or amino acid residues from at or about 37 to at or about 218 of Figure 19 (SEQ ID NO:25), or amino acid residues from at or about 12 to at or about 218 of Figure 19 (SEQ ID NO:25), or amino acid residues from at or about 1 to at or about 218 of Figure 19 (SEQ ID NO:25).

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Human interleukin-1 receptor antagonist analog 1 short ("hlL-1Ra15"), hlL-1Ra1S polypeptide, and hlL-1Ra1S protein are defined as any native sequence hlL-1Ra1S or hlL-1Ra1S variant (which are further defined herein).

A "native sequence hIL-1Ra1S" means a polypeptide comprising a naturally-occurring amino acid sequence selected from the group consisting of: (1) the amino acid sequence of amino acid residues from at or about 1 to at or about 38 of Figure 16 (SEQ ID NO:21); (2) the amino acid sequence of amino acid residues from at or about 26 to at or about 167 of Figure 16 (SEQ ID NO:21); (3) the amino acid sequence of amino acid residues from at or about 39 to at or about 167 of Figure 16 (SEQ ID NO:21); (4) the amino acid sequence of amino acid residues from at or about 47 to at or about 167 of Figure 16 (SEQ ID NO:21); and (5) the amino acid sequence of any naturally-occurring truncated or secreted form or any naturally-occurring allelic variant of a polypeptide comprising the amino acid sequence of (1) or (2) or (3) or (4). In one embodiment of the invention, the native sequence hIL-1Ra1S comprises amino acids from at or about 26 to at or about 167 of Figure 16 (SEQ ID NO:21), or amino acids from at or about 1 to at or about 167 of Figure 16 (SEQ ID NO:21). In another embodiment, the native sequence hIL-1Ra1S consists of amino acids from at or about 39 to at or about 167 of Figure 16 (SEQ ID NO:21) or amino acids from at or about 39 to at or about 167 of Figure 16 (SEQ ID NO:21).

"hIL-1Ra1S fusion variant" and "hIL-1Ra1S variant" mean a chimeric hIL-1Ra1S consisting of a native sequence hIL-1Ra1S fused at its N-terminus or C-terminus to a heterologous amino acid or amino acid sequence. In one embodiment, the hIL-1Ra1S fusion variant polypeptide consists of a native sequence of hIL-1Ra1S fused at its N-terminus or C-terminus to a heterologous amino acid or amino acid sequence, wherein the heterologous amino acid or amino acid sequence is heterologous to the native sequence, i.e. the resulting chimeric sequence is non-naturally occurring. In another embodiment, the hIL-1Ra1S fusion variant consists of the amino acid sequence of amino acids from at or about 47 to at or about 167 of Figure 16 (SEQ ID NO:21), or the amino acid sequence of amino acids from at or about 39 to at or about 167 of Figure 16 (SEQ ID NO:21), fused at its N-terminus or C-terminus to a heterologous amino acid or amino acid sequence to form a non-naturally occurring fusion protein. Such hIL-1Ra1S fusion variants include, for instance, hIL-1Ra1S polypeptides wherein a heterologous secretion leader sequence is fused to the N-terminus of the sequence of amino acid residues from at or about 47 to at or about 167 of Figure 16 (SEQ ID NO:21), or amino acid residues from at or about 39 to at or about 167 of Figure 16 (SEQ ID NO:21), or amino acid residues from at or about 39 to at or about 167 of Figure 16 (SEQ ID NO:21), or amino acid residues from at or about 39 to at or about 167 of Figure 16 (SEQ ID NO:21).

Human interleukin-1 receptor antagonist analog 2 ("hIL-1Ra2"), hIL-1Ra2 polypeptide, and hIL-1Ra2 protein are defined as any native sequence hIL-1Ra2 or hIL-1Ra2 fusion variant (which are further defined herein).

A "native sequence hIL-1Ra2" means (1) a polypeptide comprising the amino acid sequence of amino acid residues from at or about 1 to at or about 134 of Figure 5 (SEQ ID NO:10) or (2) a polypeptide consisting of a naturally-occurring truncated or secreted form of the polypeptide of (1). In one embodiment of the invention, the native sequence hIL-1Ra2 consists of amino acids from at or about 27 to at or about 134 of Figure 5 (SEQ ID NO:10), or amino acids from at or about 1 to at or about 134 of Figure 5 (SEQ ID NO:10).

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WO 00/39297 PCT/US99/30720

"hIL-1Ra2 fusion variant" and "hIL-1Ra2 variant" mean a chimeric hIL-1Ra2 consisting of a native sequence hIL-1Ra2 fused at its N-terminus or C-terminus to a heterologous amino acid or amino acid sequence. In one embodiment, the hIL-1Ra2 fusion variant polypeptide consists of a native sequence of hIL-1Ra2 fused at its N-terminus or C-terminus to a heterologous amino acid or amino acid sequence, wherein the heterologous amino acid or amino acid sequence is heterologous to the native sequence, i.e. the resulting chimeric sequence is non-naturally occurring. In another embodiment, the hIL-1Ra2 variant consists of the amino acid sequence of amino acids from at or about 27 to at or about 134 of Figure 5 (SEQ ID NO:10), or amino acids from at or about 1 to at or about 134 of Figure 5 (SEQ ID NO:10), fused at its N-terminus or C-terminus to a heterologous amino acid or amino acid sequence to form a non-naturally occurring fusion protein. Such hIL-1Ra2 fusion variants include, for instance, hIL-1Ra2 polypeptides wherein a heterologous secretion leader sequence is fused to the N-terminus of the sequence of amino acids from at or about 27 to at or about 134 of Figure 5 (SEQ ID NO:10), or amino acids from at or about 1 to at or about 134 of Figure 5 (SEQ ID NO:10), or amino acids from at or about 1 to at or about 134 of Figure 5 (SEQ ID NO:10).

Human interleukin-1 receptor antagonist analog 3 ("hIL-1Ra3"), hIL-1Ra3 polypeptide, and hIL-1Ra3 protein are defined as any native sequence hIL-1Ra3 or variant hIL-1Ra3 (which are further defined herein).

A "native sequence hIL-1Ra3" means a polypeptide comprising an amino acid sequence selected from the group consisting of: (1) the amino acid sequence of amino acid residues from at or about 95 to at or about 134 of Figure 7 (SEQ ID NO:13); (2) the amino acid sequence of amino acid residues from at or about 34 to at or about 155 of Figure 7 (SEQ ID NO:13); and (3) the amino acid sequence of any naturally-occurring truncated or secreted form or any naturally-occurring allelic variant of a polypeptide comprising the amino acid sequence of (1) or (2). In one embodiment of the invention, the native sequence hIL-1Ra3 comprises amino acids from at or about 34 to at or about 155 of Figure 7 (SEQ ID NO:13), or amino acids from at or about 2 to at or about 155 of Figure 7 (SEO ID NO:13).

"hIL-1Ra3 variant" is defined as any hIL-1Ra3 C-terminal variant or hIL-1Ra3 full sequence variant (which are further defined herein).

"hIL-1Ra3 C-terminal variant" means any hIL-1Ra3 other than a native sequence hIL-1Ra3, which variant is an active hIL-1Ra3, as defined below, having at least about 80% amino acid sequence identity with the amino acid sequence of amino acid residues from at or about 95 to at or about 134 of Figure 7 (SEQ ID NO:13) or the amino acid sequence of amino acid residues from at or about 80 to at or about 155 of Figure 7 (SEQ ID NO:13). Such hIL-1Ra3 C-terminal variants include, for instance, hIL-1Ra3 polypeptides wherein one or more amino acid residues are added, or deleted, internally or at the N- or C-terminus, in the sequence of amino acid residues from at or about 95 to at or about 134 of Figure 7 (SEQ ID NO:13) or in the sequence of amino acid residues from at or about 80 to at or about 155 of Figure 7 (SEQ ID NO:13). Ordinarily, an hIL-1Ra3 C-terminal variant will have at least about 80% amino acid sequence identity, or at least about 85% amino acid sequence identity, or at least about 85% amino acid sequence identity, or at least about 90% amino acid sequence identity, or at least about 95% amino acid sequence identity with the amino acid sequence of amino acid residues from at or about 95 to at or about 134 of Figure 7 (SEQ ID

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WO 00/39297 PCT/US99/30720

NO:13) or the amino acid sequence of amino acid residues from at or about 80 to at or about 155 of Figure 7 (SEO ID NO:13).

"hIL-1Ra3 full sequence variant" means any hIL-1Ra3 other than a native sequence hIL-1Ra3, which variant retains at least one biologic activity of a native sequence hIL-1Ra3, such as the ability to bind IL-1R, and which variant has at least about 80% amino acid sequence identity, or at least about 85% amino acid sequence identity, or at least about 95% amino acid sequence identity with the amino acid sequence of amino acid residues from at or about 34 to at or about 155 of Figure 7 (SEQ ID NO:13) or the amino acid sequence of acid residues from at or about 2 to at or about 155 of Figure 7 (SEQ ID NO:13). Such hIL-1Ra3 full sequence variants include, for instance, hIL-1Ra1 polypeptides wherein one or more amino acid residues are added, or deleted, internally or at the N- or C-terminus, in the sequence of amino acid residues from at or about 34 to at or about 155 of Figure 7 (SEQ ID NO:13) or the amino acid sequence of amino acid residues from at or about 25 to at or about 155 of Figure 7 (SEQ ID NO:13).

Murine interleukin-1 receptor antagonist analog 3 ("mIL-1Ra3"), mIL-1Ra3 polypeptide, and mIL-1Ra3 protein are defined as any native sequence mIL-1Ra3 or variant mIL-1Ra3.

A "native sequence mIL-1Ra3" means a polypeptide comprising an amino acid sequence selected from the group consisting of: (1) the amino acid sequence of amino acid residues from at or about 95 to at or about 134 of Figure 9 (SEQ ID NO:16); (2) the amino acid sequence of amino acid residues from at or about 34 to at or about 155 of Figure 9 (SEQ ID NO:16); and (3) the amino acid sequence of any naturally-occurring truncated or secreted form or naturally-occurring allelic variant of a polypeptide comprising the amino acid sequence of (1) or (2). In one embodiment of the invention, the native sequence mIL-1Ra3 comprises amino acids from at or about 34 to at or about 155 of Figure 9 (SEQ ID NO:16).

"mIL-1Ra3 variant" is defined as any mIL-1Ra3 C-terminal variant or mIL-1Ra3 full sequence variant (which are further defined herein).

"mIL-1Ra3 C-terminal variant" means any mIL-1Ra3 other than a native sequence mIL-1Ra3, which variant is an active mIL-1Ra3, as defined below, having at least about 80% amino acid sequence identity with the amino acid sequence of amino acids from at or about 95 to at or about 134 of Figure 9 (SEQ ID NO:16). Such mIL-1Ra3 C-terminal variants include, for instance, mIL-1Ra3 polypeptides wherein one or more amino acid residues are added, or deleted, internally or at the N- or C-terminus, in the sequence of amino acids from at or about 95 to at or about 134 of Figure 9 (SEQ ID NO:16). Ordinarily, an mIL-1Ra3 C-terminal variant will have at least about 80% amino acid sequence identity, or at least about 95% amino acid sequence identity, and or at least about 95% amino acid sequence identity, and or at least about 95% amino acid sequence identity, and or at least about 95% amino acid sequence identity with the amino acid sequence of amino acids 95 to 134 of Figure 9 (SEQ ID NO:16).

"mIL-1Ra3 full sequence variant" means any mIL-1Ra3 other than a native sequence mIL-1Ra3, which variant retains at least one biologic activity of a native sequence mIL-1Ra3, such as the ability to bind IL-1R, and which variant has at least about 85% amino acid sequence identity, or at least about 95% sequence

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identity with the amino acid sequence of amino acid residues from at or about 34 to at or about 155 of Figure 9 (SEQ ID NO:16) or the amino acid sequence of amino acid residues from at or about 2 to at or about 155 of Figure 9 (SEQ ID NO:16). Such mIL-1Ra3 full sequence variants include, for instance, mIL-1Ra3 polypeptides wherein one or more amino acid residues are added, or deleted, internally or at the N- or C-terminus, in the sequence of amino acid residues from at or about 34 to at or about 155 of Figure 9 (SEQ ID NO:16) or in the sequence of amino acid residues from at or about 2 to at or about 155 of Figure 9 (SEQ ID NO:16).

"Human interleukin-1-like polypeptide", "hIL-1lp", "hIL-1lp polypeptide", "hIL-1lp protein", "human interleukin-1 receptor antagonist analog", "hIL-1Raa", "hIL-1Raa polypeptide", and "hIL-1Raa protein" are defined as any hIL-1Ra1, hIL-1Ra2 or hIL-1Ra3 polypeptide.

"Native sequence hIL-1lp" and "native sequence hIL-1Raa" are defined as any polypeptide that comprises a native sequence hIL-1Ra1, hIL-1Ra2, or hIL-1Ra3.

"hIL-11p variant" is defined as any polypeptide that comprises a variant of hIL-1Ra1, hIL-1Ra2, or hIL-1Ra3.

"Interleukin-1 receptor", "interleukin-1 receptor polypeptide", "interleukin-1 receptor protein", "IL-1 receptor", "IL-1R", "IL-1R polypeptide", and "IL-1R protein", are defined as the family of cell surface proteins that bind to interleukin-1 (IL-1) and/or function in IL-1-induced signal transduction in a given species, such as human or mouse. IL-1R includes the human T cell-expressed IL-1 receptor disclosed in Sims, et al., Proc. Natl. Acad. Sci. (USA), 86: 8946-8950 (1989).

"Interleukin-18 receptor", "interleukin-18 receptor polypeptide", "interleukin-18 receptor protein", "IL-18 receptor", "IL-18R", "IL-18R polypeptide", and "IL-18R protein", are defined as the family of cell surface proteins that bind to interleukin-18 (IL-18) and/or function in IL-18induced signal transduction in a given species, such as human or mouse. IL-18R includes the IL-1 receptor related protein (IL-1Rrp) described in Torigoe et al., J. Biol. Chem., 272: 25737-25742 (1997) and the 1L-18 receptor accessory protein-like molecule (IL-18RAcPL) described in Born et al., J. Biol. Chem., 273: 29445-29450 (1998).

"Interleukin-1-like family" and "IL-1-like family" are used to indicate the family of polypeptides related to the ligands of IL-1R or IL-18R. The IL-1-like family includes IL-1 receptor agonists and antagonists and related polypeptides such as IL-1α (described in Bazan et al., Nature, 379: 591 (1996), IL-18 (Bazan et al.), IL-18 (interferon-y inducing factor)(IGIF)(Bazan et al.), IL-1 receptor antagonist polypeptides such as secretory IL-1Ra (sIL-1Ra)(described in Eisenberg et al., Nature, 343: 341-346 (1990)) and intracellular IL-1Ra (iclL-1Ra) (described in Haskill et al., Proc. Natl. Acad. Sci. (USA), 88: 3681-3685 (1991)), and the IL-11p polypeptides of the invention.

"Percent (%) amino acid sequence identity" with respect to the IL-11p sequences identified herein is defined as the percentage of amino acid residues in a candidate sequence that are identical with the amino acid residues in an IL-1lp sequence, after aligning the sequences and introducing gaps, if necessary, to achieve the maximum percent sequence identity, and not considering any conservative substitutions as part of the sequence identity. Alignment for purposes of determining percent amino acid sequence identity can be achieved in various ways

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that are within the skill in the art, for instance, using publicly available computer software such as BLAST, BLAST-2, ALIGN, ALIGN-2 or Megalign (DNASTAR) software. Those skilled in the art can determine appropriate parameters for measuring alignment, including any algorithms needed to achieve maximal alignment over the full-length of the sequences being compared. For purposes herein, however, % amino acid sequence identity values are obtained as described below by using the sequence comparison computer program ALIGN-2, wherein the complete source code for the ALIGN-2 program is provided in Tables 3A-3Q. The ALIGN-2 sequence comparison computer program was authored by Genentech, Inc. and the source code shown in Tables 3A-3Q has been filed with user documentation in the U.S. Copyright Office, Washington D.C., 20559, where it is registered under U.S. Copyright Registration No. TXU510087. The ALIGN-2 program is publicly available through Genentech, Inc., South San Francisco, California or may be compiled from the source code provided in Tables 3A-3Q. The ALIGN-2 program should be compiled for use on a UNIX operating system, preferably digital UNIX V4.0D. All sequence comparison parameters are set by the ALIGN-2 program and do not vary.

For purposes herein, the % amino acid sequence identity of a given amino acid sequence A to, with, or against a given amino acid sequence B (which also can be phrased as a given amino acid sequence A that has or comprises a certain % amino acid sequence identity to, with, or against a given amino acid sequence B) is calculated as follows:

### 100 times the fraction X/Y

where X is the number of amino acid residues scored as identical matches by the sequence alignment program ALIGN-2 in that program's alignment of A and B, and where Y is the total number of amino acid residues in B. It will be appreciated that where the length of amino acid sequence A is not equal to the length of amino acid sequence B, the % amino acid sequence identity of A to B will not equal the % amino acid sequence identity of B to A. As examples of % amino acid sequence identity calculations, Tables 2A-2B demonstrate how to calculate the % amino acid sequence identity of the amino acid sequence designated "Comparison Protein" to the amino acid sequence designated "PRO".

Unless specifically stated otherwise, all % amino acid sequence identity values used herein are obtained as described above using the ALIGN-2 sequence comparison computer program. However, % amino acid sequence identity may also be determined using the sequence comparison program NCBI-BLAST2 (Altschul et al., Nucleic Acids Res. 25:3389-3402 (1997)). The NCBI-BLAST2 sequence comparison program may be downloaded from http://www.ncbi.nlm.nih.gov. NCBI-BLAST2 uses several search parameters, wherein all of those search parameters are set to default values including, for example, unmask = yes, strand = all, expected occurrences = 10, minimum low complexity length = 15/5, multi-pass e-value = 0.01, constant for multi-pass = 25, dropoff for final gapped alignment = 25 and scoring matrix = BLOSUM62.

In situations where NCBl-BLAST2 is employed for amino acid sequence comparisons, the % amino acid sequence identity of a given amino acid sequence A to, with, or against a given

PCT/US99/30720

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amino acid sequence B (which also can be phrased as a given amino acid sequence A that has or comprises a certain % amino acid sequence identity to, with, or against a given amino acid sequence B) is calculated as follows:

100 times the fraction X/Y

where X is the number of amino acid residues scored as identical matches by the sequence alignment program NCBI-BLAST2 in that program's alignment of A and B, and where Y is the total number of amino acid residues in B. It will be appreciated that where the length of amino acid sequence A is not equal to the length of amino acid sequence B, the % amino acid sequence identity of A to B will not equal the % amino acid sequence identity of B to A.

"Percent (%) nucleic acid sequence identity" with respect to the IL-11p polypeptideencoding nucleic acid sequences identified herein is defined as the percentage of nucleotides in a candidate sequence that are identical with the nucleotides in an IL-1lp polypeptide-encoding nucleic acid sequence, after aligning the sequences and introducing gaps, if necessary, to achieve the maximum percent sequence identity. Alignment for purposes of determining percent nucleic acid sequence identity can be achieved in various ways that are within the skill in the art, for instance, using publicly available computer software such as BLAST, BLAST-2, ALIGN, ALIGN-2 or Megalign (DNASTAR) software. Those skilled in the art can determine appropriate parameters for measuring alignment, including any algorithms needed to achieve maximal alignment over the full-length of the sequences being compared. For purposes herein, however, % nucleic acid sequence identity values are obtained as described below by using the sequence comparison computer program ALIGN-2, wherein the complete source code for the ALIGN-2 program is provided in Tables 3A-3Q. The ALIGN-2 sequence comparison computer program was authored by Genentech, Inc. and the source code shown in Tables 3A-3Q has been filed with user documentation in the U.S. Copyright Office, Washington D.C., 20559, where it is registered under U.S. Copyright Registration No. TXU510087. The ALIGN-2 program is publicly available through Genentech, Inc., South San Francisco, California or may be compiled from the source code provided in Tables 3A-3Q. The ALIGN-2 program should be compiled for use on a UNIX operating system, preferably digital UNIX V4.0D. All sequence comparison parameters are set by the ALIGN-2 program and do not vary.

For purposes herein, the % nucleic acid sequence identity of a given nucleic acid sequence C to, with, or against a given nucleic acid sequence D (which also can be phrased as a given nucleic acid sequence C that has or comprises a certain % nucleic acid sequence identity to, with, or against a given nucleic acid sequence D) is calculated as follows:

### 100 times the fraction W/Z

where W is the number of nucleotides scored as identical matches by the sequence alignment program ALIGN-2 in that program's alignment of C and D, and where Z is the total number of nucleotides in D. It will be appreciated that where the length of nucleic acid sequence C is not

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equal to the length of nucleic acid sequence D, the % nucleic acid sequence identity of C to D will not equal the % nucleic acid sequence identity of D to C. As examples of % nucleic acid sequence identity calculations, Tables 2C-2D demonstrate how to calculate the % nucleic acid sequence identity of the nucleic acid sequence designated "Comparison DNA" to the nucleic acid sequence designated "PRO-DNA".

Unless specifically stated otherwise, all % nucleic acid sequence identity values used herein are obtained as described above using the ALIGN-2 sequence comparison computer program. However, % nucleic acid sequence identity may also be determined using the sequence comparison program NCBI-BLAST2 (Altschul et al., Nucleic Acids Res. 25:3389-3402 (1997)). The NCBI-BLAST2 sequence comparison program may be downloaded from http://www.ncbi.nlm.nih.gov. NCBI-BLAST2 uses several search parameters, wherein all of those search parameters are set to default values including, for example, unmask = yes, strand = all, expected occurrences = 10, minimum low complexity length = 15/5, multi-pass e-value = 0.01, constant for multi-pass = 25, dropoff for final gapped alignment = 25 and scoring matrix = BLOSUM62.

In situations where NCBI-BLAST2 is employed for sequence comparisons, the % nucleic acid sequence identity of a given nucleic acid sequence C to, with, or against a given nucleic acid sequence D (which also can be phrased as a given nucleic acid sequence C that has or comprises a certain % nucleic acid sequence identity to, with, or against a given nucleic acid sequence D) is calculated as follows:

### 100 times the fraction W/Z

where W is the number of nucleotides scored as identical matches by the sequence alignment program NCBI-BLAST2 in that program's alignment of C and D, and where Z is the total number of nucleotides in D. It will be appreciated that where the length of nucleic acid sequence C is not equal to the length of nucleic acid sequence D, the % nucleic acid sequence identity of C to D will not equal the % nucleic acid sequence identity of D to C.

The term "positives", in the context of the amino acid sequence identity comparisons performed as described above, includes amino acid residues in the sequences compared that are not only identical, but also those that have similar properties. Amino acid residues that score a positive value to an amino acid residue of interest are those that are either identical to the amino acid residue of interest or are a preferred substitution (as defined in Table 1 below) of the amino acid residue of interest.

For purposes herein, the % value of positives of a given amino acid sequence A to, with, or against a given amino acid sequence B (which also can be phrased as a given amino acid sequence A that has or comprises a certain % positives to, with, or against a given amino acid sequence B) is calculated as follows:

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where X is the number of amino acid residues scoring a positive value as defined above by the sequence alignment program ALIGN-2 in that program's alignment of A and B, and where Y is the total number of amino acid residues in B. It will be appreciated that where the length of amino acid sequence A is not equal to the length of amino acid sequence B, the % positives of A to B will not equal the % positives of B to A.

"Isolated," when used to describe the various polypeptides disclosed herein, means polypeptide that has been identified and separated and/or recovered from a component of its natural environment. Contaminant components of its natural environment are materials that would typically interfere with diagnostic or therapeutic uses for the polypeptide, and may include enzymes, hormones, and other proteinaceous or non-proteinaceous solutes. In preferred embodiments, the polypeptide will be purified (1) to a degree sufficient to obtain at least 15 residues of N-terminal or internal amino acid sequence by use of a spinning cup sequenator, or (2) to homogeneity by SDS-PAGE under non-reducing or reducing conditions using Coomassie blue or, preferably, silver stain. Isolated polypeptide includes polypeptide in situ within recombinant cells, since at least one component of the IL-1lp natural environment will not be present. Ordinarily, however, isolated polypeptide will be prepared by at least one purification step.

An "isolated" nucleic acid molecule encoding a IL-11p polypeptide is a nucleic acid molecule that is identified and separated from at least one contaminant nucleic acid molecule with which it is ordinarily associated in the natural source of the IL-11p-encoding nucleic acid. An isolated IL-11p-encoding nucleic acid molecule is other than in the form or setting in which it is found in nature. Isolated nucleic acid molecules therefore are distinguished from the IL-11p-encoding nucleic acid molecule as it exists in natural cells. However, an isolated nucleic acid molecule encoding a IL-11p polypeptide includes IL-11p-encoding nucleic acid molecules contained in cells that ordinarily express IL-11p where, for example, the nucleic acid molecule is in a chromosomal location different from that of natural cells.

The term "control sequences" refers to DNA sequences necessary for the expression of an operably linked coding sequence in a particular host organism. The control sequences that are suitable for prokaryotes, for example, include a promoter, optionally an operator sequence, and a ribosome binding site. Eukaryotic cells are known to utilize promoters, polyadenylation signals, and enhancers.

Nucleic acid is "operably linked" when it is placed into a functional relationship with another nucleic acid sequence. For example, DNA for a presequence or secretory leader is operably linked to DNA for a polypeptide if it is expressed as a preprotein that participates in the secretion of the polypeptide; a promoter or enhancer is operably linked to a coding sequence if it affects the transcription of the sequence; or a ribosome binding site is operably linked to a coding sequence if it is positioned so as to facilitate translation. Generally, "operably linked" means that the DNA sequences being linked are contiguous, and, in the case of a secretory leader, contiguous and in reading phase. However, enhancers do not have to be contiguous. Linking is accomplished by ligation at convenient restriction sites. If such sites do not exist,

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the synthetic oligonucleotide adaptors or linkers are used in accordance with conventional practice.

The term "antibody" is used in the broadest sense and specifically covers single anti-IL-11p monoclonal antibodies (including agonist, antagonist, and neutralizing antibodies) and anti-11L-11p antibody compositions with polyepitopic specificity. The term "monoclonal antibody" as used herein refers to an antibody obtained from a population of substantially homogeneous antibodies, i.e., the individual antibodies comprising the population are identical except for possible naturally-occurring mutations that may be present in minor amounts.

"Stringency" of hybridization reactions is readily determinable by one of ordinary skill in the art, and generally is an empirical calculation dependent upon probe length, washing temperature, and salt concentration. In general, longer probes require higher temperatures for proper annealing, while shorter probes need lower temperatures. Hybridization generally depends on the ability of denatured DNA to reanneal when complementary strands are present in an environment below their melting temperature. The higher the degree of desired homology between the probe and hybridizable sequence, the higher the relative temperature which can be used. As a result, it follows that higher relative temperatures would tend to make the reaction conditions more stringent, while lower temperatures less so. For additional details and explanation of stringency of hybridization reactions, see Ausubel et al., Current Protocols in Molecular Biology, Wiley Interscience Publishers, (1995).

"Stringent conditions" or "high stringency conditions", as defined herein, may be identified by those that: (1) employ low ionic strength and high temperature for washing, for example 0.015 M sodium chloride/0.0015 M sodium citrate/0.1% sodium dodecyl sulfate at 50°C; (2) employ during hybridization a denaturing agent, such as formamide, for example, 50% (v/v) formamide with 0.1% bovine serum albumin/0.1% Ficoil/0.1% polyvinylpyrrolidone/50mM sodium phosphate buffer at pH 6.5 with 750 mM sodium chloride, 75 mM sodium citrate at 42°C; or (3) employ 50% formamide, 5 x SSC (0.75 M NaCl, 0.075 M sodium citrate), 50 mM sodium phosphate (pH 6.8), 0.1% sodium pyrophosphate, 5 x Denhardt's solution, sonicated salmon sperm DNA (50 μg/ml), 0.1% SDS, and 10% dextran sulfate at 42°C, with washes at 42°C in 0.2 x SSC (sodium chloride/sodium citrate) and 50% formamide at 55°C, followed by a high-stringency wash consisting of 0.1 x SSC containing EDTA at 55°C.

"Moderately stringent conditions" may be identified as described by Sambrook et al., Molecular Cloning: A Laboratory Manual, New York: Cold Spring Harbor Press, 1989, and include the use of washing solution and hybridization conditions (e.g., temperature, ionic strength aind %SDS) less stringent that those described above. An example of moderately stringent conditions is overnight incubation at  $37^{\circ}$ C in a solution comprising: 20% formamide, 5 x SSC (150 mM NaCl, 15 mM trisodium citrate), 50 mM sodium phosphate (pH 7.6), 5 x Denhardt's solution, 10% dextran sulfate, and 20 mg/mL denatured sheared salmon sperm DNA, followed by washing the filters in 1 x SSC at about  $37\text{-}50^{\circ}$ C. The skilled artisan will recognize how to adjust the temperature, ionic strength, etc. as necessary to accommodate factors such as probe length and the like.

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The term "epitope tagged" when used herein refers to a chimeric polypeptide comprising an IL-11p polypeptide fused to a "tag polypeptide". The tag polypeptide has enough residues to provide an epitope against which an antibody can be made, yet is short enough such that it does not interfere with activity of the polypeptide to which it is fused. The tag polypeptide preferably also is fairly unique so that the antibody does not substantially cross-react with other epitopes. Suitable tag polypeptides generally have at least six amino acid residues and usually between about 8 and 50 amino acid residues (preferably, between about 10 and 20 amino acid residues).

As used herein, the term "immunoadhesin" designates antibody-like molecules which combine the binding specificity of a heterologous protein (an "adhesin") with the effector functions of immunoglobulin constant domains. Structurally, the immunoadhesins comprise a fusion of an amino acid sequence with the desired binding specificity which is other than the antigen recognition and binding site of an antibody (i.e., is "heterologous"), and an immunoglobulin constant domain sequence. The adhesin part of an immunoadhesin molecule typically is a contiguous amino acid sequence comprising at least the binding site of a receptor or a ligand. The immunoglobulin constant domain sequence in the immunoadhesin may be obtained from any immunoglobulin, such as IgG-1, IgG-2, IgG-3, or igG-4 subtypes, IgA (including IgA-1 and IgA-2), IgE, IgD or IgM.

"Active" or "activity" for the purposes herein refers to form(s) of IL-11p which retain one or more of the biologic activities of native or naturally-occurring IL-11p, or which exhibit immunological cross-reactivity with a native or naturally-occurring IL-11p.

As used herein, a "biologic activity" or "biological activity" of an IL-1lp means any effector function exhibited by the IL-1lp in the physiology or pathophysiology of a mammal, excluding any immunogenic or antigenic functions of the IL-1lp. Immunogenic and antigenic functions of an IL-1lp refer to the ability of the IL-1lp to generate a humoral or cell-mediated immune response specific to the IL-1lp, and the ability of the IL-1lp to specifically recognize and interact with anti-IL-Ilp antibodies. B cells or T cells, respectively, in a mammal.

As used herein, "immunological cross-reactivity" with an IL-11p means that the candidate polypeptide is capable of competitively inhibiting the binding of the IL-11p to polyclonal or monoclonal antibodies raised against the IL-11p.

In one embodiment, IL-1lp activity includes the ability to agonize or antagonize one or more biological activities of any IL-1-like family member, e.g. an IL-1lp activity that antagonizes an IL-1-mediated or IL-18-mediated inflammatory response. In another embodiment, IL-1lp activity includes the ability to bind to the IL-18 receptor and/or IL-1 receptor.

The term "antagonist" is used in the broadest sense, and includes any molecule that partially or fully blocks, inhibits, or neutralizes a biological activity of a native IL-1lp polypeptide disclosed herein. In a similar manner, the term "agonist" is used in the broadest sense and includes any molecule that mimics a biological activity of a native IL-1lp polypeptide disclosed herein. Suitable agonist or antagonist molecules specifically include agonist or antagonist antibodies or antibody fragments, fragments or amino acid sequence variants of native IL-1lp polypeptides, peptides, small organic molecules, etc.

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"Treatment" refers to both therapeutic treatment and prophylactic or preventative measures, wherein the object is to prevent or slow down (lessen) the targeted pathologic condition or disorder. Those in need of treatment include those already with the disorder as well as those prone to have the disorder or those in whom the disorder is to be prevented.

"Chronic" administration refers to administration of the agent(s) in a continuous mode as opposed to an acute mode, so as to maintain the initial therapeutic effect (activity) for an extended period of time. "Intermittent" administration is treatment that is not consecutively done without interruption, but rather is cyclic in nature.

"Mammal" for purposes of treatment refers to any animal classified as a mammal, including humans, domestic and farm animals, and zoo, sports, or pet animals, such as dogs, cats, cattle, horses, sheep, pigs, etc. Preferably, the mammal is human.

Administration "in combination with" one or more further therapeutic agents includes simultaneous (concurrent) and consecutive administration in any order.

The terms "inflammatory disorders" and "inflammatory diseases" are used interchangeably herein and refer to pathological states resulting in inflammation. Examples of such disorders include inflammatory skin diseases such as psoriasis and atopic dermatitis; systemic scleroderma and sclerosis; responses associated with inflammatory bowel disease (such as Crohn's disease and ulcerative colitis); ischemic reperfusion disorders including surgical tissue reperfusion injury, myocardial ischemic conditions such as myocardial infarction, cardiac arrest, reperfusion after cardiac surgery and constriction after percutaneous transluminal coronary angioplasty, stroke, and abdominal aortic aneurysms; cerebral edema secondary to stroke; cranial trauma; hypovolemic shock; asphyxia; adult respiratory distress syndrome; acute lung injury; Behcet's Disease; dermatomyositis; polymyositis; multiple sclerosis; dermatitis; meningitis; encephalitis; uveitis; osteoarthritis; autoimmune diseases such as rheumatoid arthritis, Sjorgen's syndrome, vasculitis, and insulin-dependent diabetes mellitus (IDDM); diseases involving leukocyte diapedesis; central nervous system (CNS) inflammatory disorder; meningitis; multiple organ injury syndrome secondary to septicaemia or trauma; inflammatory diseases of the liver, including alcoholic hepatitis and hepatic fibrosis; pathologic host responses to infection, including pathologic inflammation in granulomatous diseases, hepatitis, and bacterial pneumonia; antigen-antibody complex mediated diseases including glomerulonephritis; sepsis; sarcoidosis; immunopathologic responses to tissue/organ transplantation, including graft-versus host disease (GVHD); inflammations of the lung, including pleurisy, alveolitis, vasculitis, pneumonia, chronic bronchitis, bronchiectasis, diffuse panbronchiolitis, hypersensitivity pneumonitis, idiopathic pulmonary fibrosis (IPF), and cystic fibrosis; inflammation in renal diseases, including acute or chronic nephritic conditions such as lupus nephritis; pancreatitis; etc. The preferred indications include rheumatoid arthritis, osteoarthritis, sepsis, acute lung injury, adult respiratory distress syndrome, idiopathic pulmonary fibrosis, ischemic reperfusion (including surgical tissue reperfusion injury, stroke, myocardial ischemia, and acute myocardial infarction), asthma, psoriasis, graft-versus-host disease (GVHD), and inflammatory bowel disease such as ulcerative colitis.

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As used herein, the terms "asthma", "asthmatic disorder", "asthmatic disease", and "bronchial asthma" refer to a condition of the lungs in which there is widespread narrowing of lower airways. "Atopic asthma" and "allergic asthma" refer to asthma that is a manifestation of an IgE-mediated hypersensitivity reaction in the lower airways, including, e.g., moderate or severe chronic asthma, such as conditions requiring the frequent or constant use of inhaled or systemic steroids to control the asthma symptoms. A preferred indication is allergic asthma.

## II. Detailed Description of the Invention

The present invention provides newly identified and isolated nucleotide sequences encoding polypeptides referred to in the present application as IL-11p. In particular, cDNAs encoding IL-11p polypeptides have been identified and isolated, as disclosed in further detail in the Examples below.

Using NCBI-BLAST2 sequence alignment computer programs, it has been found that a full-length native sequence hIL-1Ra1 (shown in Figure 3 and SEQ ID NO:7) has some amino acid sequence identity with human IL-1 receptor antagonist beta (hIL-1Raß) and TANGO-77 protein, a full-length native sequence hIL-1Ra1L (shown in Figure 15 and SEQ ID NO:19) has some amino acid sequence identity with human IL-1 receptor antagonist beta (hIL-1Raß) and TANGO-77 protein, a full-length native sequence hIL-1Ra1V (shown in Figure 19 and SEQ ID NO:25) has some amino acid sequence identity with human IL-1 receptor antagonist beta (hIL-1Raß) and TANGO-77 protein, a full-length native sequence hIL-1Ra1S (shown in Figure 16 and SEQ ID NO:21) appears to be an allelic variant of TANGO-77 protein and has some amino acid sequence identity with human IL-1 receptor antagonist beta (hIL-1Raß), a full-length native sequence hIL-1Ra2 (shown in Figure 5 and SEQ ID NO:10) has some amino acid sequence identity with hIL- $1Ra\beta$ , a full-length native sequence hIL-1Ra3 (shown in Figure 7 and SEQ ID NO:13) has some amino acid sequence identity with human intracellular IL-1 receptor antagonist (hicIL-1Ra), and a full-length native sequence mIL-1Ra3 (shown in Figure 9 and SEQ ID NO:16) has some amino acid sequence identity with mouse IL-1 receptor antagonist (mIL-1Ra) and has some amino acid sequence identity with hiclL-1Ra. hlL-1Raß is described in EP 0855404 published July 29, 1998. TANGO-77 protein is described in WO 99/06426 published February 11, 1999. hicIL-1Ra is described in WO 95/10298 published April 20, 1995 and in Haskill et al., Proc. Natl. Acad. Sci. (USA), 88: 3681-3685 (1991). mIL-1Ra is described in Zahedi et al., J. Immunol., 146: 4228-4233 (1991), Matsushime et al., Blood, 78: 616-623 (1991), Zahedi et al., Cytokine, 6: 1-9 (1994), Eisenberg et al., Proc. Natl. Acad. Sci. (USA), 88: 5232-5236 (1991) and Shuck et al., Eur. J. Immunol., 21: 2775-2780 (1991). Accordingly, it is presently believed that the IL-11p polypeptides disclosed in the present application are newly identified members of the interleukin-1-like family and possess inflammatory or anti-inflammatory activities, or other cellular response activating or inhibiting activities, typical of the IL-1-like family.

In addition to the full-length native sequence IL-1lp polypeptides described herein, it is contemplated that IL-1lp variants can be prepared. Such embodiments of the invention include all IL-1lp polypeptides that are IL-1lp variants as defined herein, such as hIL-1Ra1 variants, hIL-1Ra1L variants, hIL-1Ra1S variants, hIL-1Ra2 variants, hIL-1Ra3 variants, hIL-1Ra3 variants.

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IL-1Ip variants can be prepared by introducing appropriate nucleotide changes into the IL-1Ip DNA, and/or by synthesis of the desired IL-1Ip polypeptide. Those skilled in the art will appreciate that amino acid changes may alter post-translational processes of the IL-1Ip, such as changing the number or position of glycosylation sites or altering the membrane anchoring characteristics.

Variations in the native full-length sequence IL-1lp or in various domains of the IL-1lp described herein, can be made, for example, using any of the techniques and guidelines for conservative and non-conservative mutations set forth, for instance, in U.S. Patent No. 5,364,934. Variations may be a substitution, deletion or insertion of one or more codons encoding the IL-1lp that results in a change in the amino acid sequence of the IL-1lp as compared with the native sequence IL-1lp. Optionally the variation is by substitution of at least one amino acid with any other amino acid in one or more of the domains of the IL-1lp. Guidance in determining which amino acid residue may be inserted, substituted or deleted without adversely affecting the desired activity may be found by comparing the sequence of the IL-1lp with that of homologous known protein molecules and minimizing the number of amino acid sequence changes made in regions of high homology. Amino acid substitutions can be the result of replacing one amino acid with another amino acid having similar structural and/or chemical properties, such as the replacement of a leucine with a serine, i.e., conservative amino acid replacements. Insertions or deletions may optionally be in the range of 1 to 5 amino acids. The variation allowed may be determined by systematically making insertions, deletions or substitutions of amino acids in the sequence and testing the resulting variants for activity in the in vitro assay described in the Examples below.

Table 1 below lists conservative amino acid substitutions (under the heading of "Preferred Substitutions") that are useful in generating variants of the native sequence IL-Ilp. If such substitutions result in alteration of biological activity, it is useful to introduce more substantial changes, such as the "Exemplary Substitutions" denoted in Table 1 or the substantial changes described below in reference to amino acid classes, at the active site in question.

	Original	Exemplary	Preferred
	Residue	Substitutions	Substitutions
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	Ala (A)	val; leu; ile	val
	Arg (R)	lys; gln; asn	lys
	Asn (N)	gln; his; lys; arg	gln
	Asp (D)	glu	glu
10	Cys (C)	ser	ser
	Gln (Q)	asn	asn
	Glu (E)	asp	asp
	Gly (G)	pro; ala	ala
	His (H)	asn; gln; lys; arg	arg
15	Ile (I)	leu; val; met; ala; phe;	
		norleucine	leu
	Leu (L)	norleucine; ile; val;	
		met; ala; phe	ile
	Lys (K)	arg; gln; asn	arg
20	Met (M)	leu; phe; ile	leu
	Phe (F)	leu; val; ile; ala; tyr	leu
	Pro (P)	ala	ala
**	Ser (S)	thr	thr
3	Thr (T)	ser	ser
25	Trp (W)	tyr; phe	tyr
9	Tyr (Y)	trp; phe; thr; ser	phe
1 0 0 25 1	Val (V)	ile; leu; met; phe;	•
1		ala; norleucine	leu

Substantial modifications in function or immunological identity of the IL-11p polypeptide are accomplished by selecting substitutions that differ significantly in their effect on maintaining (a) the structure of the polypeptide backbone in the area of the substitution, for example, as a sheet or helical conformation, (b) the charge or hydrophobicity of the molecule at the target site, or (c) the bulk of the side chain. Naturally occurring residues are divided into groups based on common side-chain properties:

- (1) hydrophobic: norleucine, met, ala, val, leu, ile;
- (2) neutral hydrophilic: cvs, ser, thr;
- (3) acidic: asp, glu;
- (4) basic: asn, gln, his, lys, arg;
- 40 (5) residues that influence chain orientation: gly, pro; and
  - (6) aromatic: trp, tyr, phe.

Non-conservative substitutions will entail exchanging a member of one of these classes for another class. Such substituted residues also may be introduced into the conservative substitution sites or, more preferably, into the remaining (non-conserved) sites.

The variations can be made using methods known in the art such as oligonucleotidemediated (site-directed) mutagenesis, alanine scanning, and PCR mutagenesis. Site-directed mutagenesis (Carter et al., Nucl. Acids Res., 13:4331 (1986); Zoller et al., Nucl. Acids Res., 10:6487 (1987)), cassette mutagenesis (Wells et al., Gene, 34:315 (1985)), restriction selection

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mutagenesis [Wells et al., Philos. Trans. R. Soc. London SerA, 317:415 (1986)] or other known techniques can be performed on the cloned DNA to produce the IL-11p variant DNA.

Scanning amino acid analysis can also be employed to identify one or more amino acids along a contiguous sequence. Among the preferred scanning amino acids are relatively small, neutral amino acids. Such amino acids include alanine, glycine, serine, and cysteine. Alanine is typically a preferred scanning amino acid among this group because it eliminates the sidechain beyond the beta-carbon and is less likely to alter the main-chain conformation of the variant [Cunningham and Wells, Science, 244: 1081-1085 (1989)]. Alanine is also typically preferred because it is the most common amino acid. Further, it is frequently found in both buried and exposed positions [Creighton, The Proteins, (W.H. Freeman & Co., N.Y.); Chothia, J. Mol. Biol., 150:1 (1976)]. If alanine substitution does not yield adequate amounts of variant, an isoteric amino acid can be used.

Covalent modifications of IL-1lp are included within the scope of this invention. One type of covalent modification includes reacting targeted amino acid residues of an IL-1lp polypeptide with an organic derivatizing agent that is capable of reacting with selected side chains or the N-or C- terminal residues of the IL-1lp. Derivatization with bifunctional agents is useful, for instance, for crosslinking IL-1lp to a water-insoluble support matrix or surface for use in the method for purifying anti-IL-1lp antibodies, and vice-versa. Commonly used crosslinking agents include, e.g., 1,1-bis(diazoacetyl)-2-phenylethane, glutaraldehyde, N-hydroxysuccinimide esters, for example, esters with 4-azidosalicylic acid, homobifunctional inidoesters, including disuccinimidyl esters such as 3,3'-dithiobis(succinimidylpropionate), bifunctional maleimides such as bis-N-maleimido-1,8-octane and agents such as methyl-3-[(p-azidophenyl]dithio]propiomidate.

Other modifications include deamidation of glutaminyl and asparaginyl residues to the corresponding glutamyl and aspartyl residues, respectively, hydroxylation of proline and lysine, phosphorylation of hydroxyl groups of seryl or threonyl residues, methylation of the  $\alpha$ -amino groups of lysine, arginine, and histidine side chains [T.E. Creighton, *Proteins: Structure and Molecular Properties*, W.H. Freeman & Co., San Francisco, pp. 79-86 (1983)], acetylation of the N-terminal amine, and amidation of any C-terminal carboxyl group.

Another type of covalent modification of the IL-11p polypeptide included within the scope of this invention comprises altering the native glycosylation pattern of the polypeptide. "Altering the native glycosylation pattern" is intended for purposes herein to mean deleting one or more carbohydrate moieties found in native sequence IL-11p (either by removing the underlying glycosylation site or by deleting the glycosylation behenical and/or enzymatic means), and/or adding one or more glycosylation sites that are not present in the native sequence IL-11p. In addition, the phrase includes qualitative changes in the glycosylation of the native proteins, involving a change in the nature and proportions of the various carbohydrate moieties present.

Addition of glycosylation sites to the IL-11p polypeptide may be accomplished by altering the amino acid sequence. The alteration may be made, for example, by the addition of, or substitution by, one or more serine or threonine residues to the native sequence IL-11p (for O-

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linked glycosylation sites). The IL-1lp amino acid sequence may optionally be altered through changes at the DNA level, particularly by mutating the DNA encoding the IL-1lp polypeptide at preselected bases such that codons are generated that will translate into the desired amino acids.

Another means of increasing the number of carbohydrate moieties on the IL-11p polypeptide is by chemical or enzymatic coupling of glycosides to the polypeptide. Such methods are described in the art, e.g., in WO 87/05330 published 11 September 1987, and in Aplin and Wriston, CRC Crit. Rev. Biochem., pp. 259-306 (1981).

Removal of carbohydrate moieties present on the IL-1lp polypeptide may be accomplished chemically or enzymatically or by mutational substitution of codons encoding for amino acid residues that serve as targets for glycosylation. Chemical deglycosylation techniques are known in the art and described, for instance, by Hakimuddin, et al., Arch. Biochem. Biophys., 259:52 (1987) and by Edge et al., Anal. Biochem., 118:131 (1981). Enzymatic cleavage of carbohydrate moieties on polypeptides can be achieved by the use of a variety of endo- and exo-glycosidases as described by Thotakura et al., Meth. Enzymol., 138:350 (1987).

Another type of covalent modification of IL-1lp comprises linking the IL-1lp polypeptide to one of a variety of nonproteinaceous polymers, e.g., polyethylene glycol (PEG), polypropylene glycol, or polyoxyalkylenes, in the manner set forth in U.S. Patent Nos. 4,640,835; 4,496,689; 4,301,144; 4,670,417; 4,791,192 or 4,179,337.

The IL-1lp of the present invention may also be modified in a way to form a chimeric molecule comprising IL-1lp fused to another, heterologous polypeptide or amino acid sequence.

In one embodiment, such a chimeric molecule comprises a fusion of the IL-1lp with a tag polypeptide which provides an epitope to which an anti-tag antibody can selectively bind. The epitope tag is generally placed at the amino- or carboxyl- terminus of the IL-1lp. The presence of such epitope-tagged forms of the IL-11p can be detected using an antibody against the tag polypeptide. Also, provision of the epitope tag enables the IL-1lp to be readily purified by affinity purification using an anti-tag antibody or another type of affinity matrix that binds to the epitope tag. Various tag polypeptides and their respective antibodies are well known in the art. Examples include poly-histidine (poly-his) or poly-histidine-glycine (poly-his-gly) tags; the flu HA tag polypeptide and its antibody 12CA5 [Field et al., Mol. Cell. Biol., 8:2159-2165 (1988)]; the cmyc tag and the 8F9, 3C7, 6E10, G4, B7 and 9E10 antibodies thereto [Evan et al., Molecular and Cellular Biology, 5:3610-3616 (1985)]; and the Herpes Simplex virus glycoprotein D (gD) tag and its antibody [Paborsky et al., Protein Engineering, 3(6):547-553 (1990)]. Other tag polypeptides include the Flag-peptide [Hopp et al., BioTechnology, 6:1204-1210 (1988)]; the KT3 epitope peptide [Martin et al., Science, 255:192-194 (1992)]; an α-tubulin epitope peptide [Skinner et al., J. Biol. Chem., 266:15163-15166 (1991)]; and the T7 gene 10 protein peptide tag [Lutz-Frevermuth et al., Proc. Natl. Acad. Sci. USA, 87:6393-6397 (1990)].

In an alternative embodiment, the chimeric molecule may comprise a fusion of the IL-11p with an immunoglobulin or a particular region of an immunoglobulin. For a bivalent form of the chimeric molecule (also referred to as an immunoadhesin), such a fusion could be to the Fc

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region of an IgG molecule. The Ig fusions preferably include the substitution of a soluble form of an IL-11p polypeptide in place of at least one variable region within an Ig molecule. In a particularly preferred embodiment, the immunoglobulin fusion includes the hinge, CH2 and CH3, or the hinge, CH1, CH2 and CH3 regions of an IgG1 molecule. For the production of immunoglobulin fusions see also US Patent No. 5,428,130 issued June 27, 1995.

In one aspect, the invention provides an isolated nucleic acid comprising DNA encoding an IL-1lp polypeptide that retains at least one biologic activity of a native sequence IL-1lp, such as the IL-1R binding activity of a native sequence hIL-1Ra3 or mIL-1Ra3, or the IL-18R binding activity of a native sequence hIL-1Ra1, hIL-1Ra1L, or hIL-1Ra1V, which DNA has at least at or about 80% sequence identity, or at least at or about 85% sequence identity, or at least at or about 90% sequence identity, or at least at or about 95% sequence identity to (a) a DNA molecule selected from the group consisting of: (1) a DNA molecule encoding an IL-1lp polypeptide comprising amino acid residues from at or about 37 to at or about 203, inclusive of Figure 2 (SEQ ID NO:5), (2) a DNA molecule encoding an IL-1lp polypeptide comprising amino acid residues from at or about 15 to at or about 193, inclusive of Figure 3 (SEQ ID NO:7), (3) a DNA molecule encoding an lL-1lp polypeptide comprising amino acid residues from at or about 34 to at or about 155, inclusive of Figure 7 (SEQ ID NO:13), (4) a DNA molecule encoding an IL-1lp polypeptide comprising amino acid residues from at or about 34 to at or about 155, inclusive of Figure 9 (SEO ID NO:16), (5) a DNA molecule encoding an IL-11p polypeptide comprising amino acid residues from at or about 26 to at or about 207, inclusive of Figure 15 (SEQ ID NO:19), and (6) a DNA molecule encoding an IL-11p polypeptide comprising amino acid residues from at or about 46 to at or about 218 of Figure 19 (SEQ ID NO:25), or (b) the complement of the DNA molecule of (a).

In another aspect, the invention provides an isolated nucleic acid comprising DNA encoding an IL-11p polypeptide that retains at least one biologic activity of a native sequence IL-11p, such as the IL-18R binding activity of a native sequence hIL-1Ra1, hIL-1Ra1L, or hIL-1Ra1V, which DNA has at least at or about 80% sequence identity, or at least at or about 85% sequence identity, or at least at or about 90% sequence identity, or at least at or about 95% sequence identity to (a) a DNA molecule selected from the group consisting of: (1) a DNA molecule encoding an IL-11p polypeptide comprising amino acid residues from at or about 1 to at or about 207, inclusive of Figure 15 (SEQ ID NO:19), and (2) a DNA molecule encoding an IL-11p polypeptide comprising amino acid residues from at or about 1 to at or about 218 of Figure 19 (SEQ ID NO:25), or (b) the complement of the DNA molecule of (a).

In another aspect, the invention provides an isolated nucleic acid comprising DNA encoding an IL-1lp polypeptide that retains at least one biologic activity of a native sequence IL-1lp, such as the IL-18R binding activity of a native sequence hIL-1Ra1, or the IL-1R binding activity of a native sequence hIL-1Ra3, which DNA has at least at or about 80% sequence identity, or at least at or about 85% sequence identity, or at least at or about 90% sequence identity, or at least at or about 95% sequence identity to (a) a DNA molecule selected from the group consisting of: (1) a DNA molecule encoding an IL-1lp polypeptide comprising amino acid residues from at or about 95 to at or about 134, inclusive of Figure 7 (SEQ ID NO:13).

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and (2) a DNA molecule encoding an IL-11p polypeptide comprising amino acid residues from at or about 95 to at or about 134, inclusive of Figure 9 (SEQ ID NO:16), or (b) the complement of the DNA molecule of (a).

PCT/US99/30720

In another aspect, the invention provides an isolated nucleic acid comprising DNA encoding an IL-1lp polypeptide that retains at least one biologic activity of a native sequence IL-1lp, such as the IL-18R binding activity of a native sequence hlL-1Ra1, or the IL-1R binding activity of a native sequence hlL-1Ra3 or mIL-1Ra3, which DNA has at least at or about 80% sequence identity, or at least at or about 85% sequence identity, or at least at or about 90% sequence identity to (a) a DNA molecule encoding an IL-1lp polypeptide comprising amino acid residues from at or about 80 to at or about 155, inclusive of Figure 7 (SEQ ID NO:13), or (b) the complement of the DNA molecule of (a).

In another aspect, the invention provides an isolated nucleic acid comprising DNA encoding an IL-1lp polypeptide that retains at least one biologic activity of a native sequence IL-1lp, such as the IL-18R binding activity of a native sequence hIL-1Ra1, or the IL-1R binding activity of a native sequence hIL-1Ra3 or mIL-1Ra3, which DNA has at least at or about 80% sequence identity, or at least at or about 85% sequence identity, or at least at or about 90% sequence identity, or at least at or about 95% sequence identity to (a) a DNA molecule selected from the group consisting of: (1) a DNA molecule encoding an IL-1lp polypeptide comprising amino acid residues from at or about 2 to at or about 25, inclusive of Figure 7 (SEQ ID NO:13), and (2) a DNA molecule encoding an IL-1lp polypeptide comprising amino acid residues from at or about 2 to at or about 155, inclusive of Figure 9 (SEQ ID NO:16), or (b) the complement of the DNA molecule of (a).

In another aspect, the invention provides an isolated nucleic acid comprising DNA having at least at or about 80% sequence identity, or at least at or about 85% sequence identity, or at least at or about 95% sequence identity to (a) a least at or about 90% sequence identity, or at least at or about 95% sequence identity to (a) a DNA molecule selected from the group consisting of: (1) a DNA molecule encoding an IL-11p polypeptide comprising amino acid residues from at or about 95 to at or about 134, inclusive of Figure 7 (SEQ ID NO:13), and (2) a DNA molecule encoding an IL-11p polypeptide comprising amino acid residues from at or about 95 to at or about 134, inclusive of Figure 9 (SEQ ID NO:16), or (b) the complement of the DNA molecule of (a).

In another aspect, the invention provides an isolated nucleic acid comprising DNA having at least at or about 80% sequence identity, or at least at or about 85% sequence identity, or at least at or about 95% sequence identity to (a) a DNA molecule encoding an IL-Ilp polypeptide comprising amino acid residues from at or about 80 to at or about 155, inclusive of Figure 7 (SEQ ID NO:13), or (b) the complement of the DNA molecule of (a).

In another aspect, the invention provides an isolated nucleic acid comprising DNA having at least at or about 80% sequence identity, or at least at or about 85% sequence identity, or at least at or about 95% sequence identity to (a) a DNA molecule selected from the group consisting of: (1) a DNA molecule encoding an IL-11p polypeptide comprising amino acid residues from at or about 2 to at or about 155, inclusive of

WO 00/39297 PCT/US99/30720

Figure 7 (SEQ ID NO:13), and (2) a DNA molecule encoding an IL-11p polypeptide comprising amino acid residues from at or about 2 to at or about 155, inclusive of Figure 9 (SEQ ID NO:16), or (b) the complement of the DNA molecule of (a).

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In another aspect, the invention concerns an isolated nucleic acid molecule encoding an IL-11p polypeptide, comprising DNA hybridizing to the complement of a nucleic acid sequence selected from the group consisting of: (1) the nucleic acid sequence consisting of nucleotide positions from at or about 238 to at or about 465 in the sense strand of Figure 7 (SEQ ID NO:12); (2) the nucleic acid sequence consisting of nucleotide positions from at or about 427 to at or about 609 in the sense strand of Figure 9 (SEQ ID NO:15); and (3) the nucleic acid sequence consisting of nucleotide positions from at or about 79 to at or about 135 in the sense strand of Figure 15 (SEQ ID NO:18). Preferably, hybridization occurs under stringent hybridization and wash conditions.

In another aspect, the invention concerns an isolated nucleic acid molecule, comprising DNA that is at least 90 nucleotides in length and that hybridizes to the complement of a nucleic acid sequence selected from the group consisting of: (1) the nucleic acid sequence consisting of nucleotide positions from at or about 238 to at or about 465 in the sense strand of Figure 7 (SEQ ID NO:12); (2) the nucleic acid sequence consisting of nucleotide positions from at or about 427 to at or about 609 in the sense strand of Figure 9 (SEQ ID NO:15); and (3) the nucleic acid sequence consisting of nucleotide positions from at or about 115 to at or about 135 in the sense strand of Figure 15 (SEQ ID NO:18). Preferably, hybridization occurs under stringent hybridization and wash conditions.

In another aspect, the invention concerns an isolated nucleic acid molecule comprising DNA encoding an IL-1lp polypeptide that retains at least one biologic activity of a native sequence IL-1lp, such as the IL-1R binding activity of a native sequence hIL-1Ra1, or hIL-1Ra1V, which DNA hybridizes to the complement of a nucleic acid sequence selected from the group consisting of: (1) the nucleic acid sequence consisting of nucleotide positions from at or about 118 to at or about 231 in the sense strand of Figure 2 (SEQ ID NO:4); (2) the nucleic acid sequence consisting of nucleotide positions from at or about 100 to at or about 465 in the sense strand of Figure 7 (SEQ ID NO:12); (3) the nucleic acid sequence consisting of nucleotide positions from at or about 244 to at or about 609 in the sense strand of Figure 9 (SEQ ID NO:15); and (4) the nucleic acid sequence consisting of nucleotide positions from at or about 208 to at or about 339 in the sense strand of Figure 19 (SEQ ID NO:24). Preferably, hybridization occurs under stringent hybridization and wash conditions.

In another aspect, the invention concerns an isolated nucleic acid molecule comprising DNA encoding an IL-1lp polypeptide that retains at least one biologic activity of a native sequence IL-1lp, such as the IL-1R binding activity of a native sequence hIL-1Ra3 or mIL-1Ra3, which DNA hybridizes to the complement of a nucleic acid sequence selected from the group consisting of (1) the nucleic acid sequence consisting of fucleotide positions from at or about 4 to at or about 465 in the sense strand of Figure 7 (SEQ ID NO:12); and (2) the nucleic acid sequence consisting of nucleotide positions from at or about 148 to at or about 609 in the sense strand

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of Figure 9 (SEQ ID NO:15). Preferably, hybridization occurs under stringent hybridization and wash conditions.

In a further aspect, the invention concerns an isolated nucleic acid molecule comprising DNA encoding an IL-11p polypeptide that retains at least one biologic activity of a native sequence IL-1lp, such as the IL-1R binding activity of a native sequence hIL-1Ra3 or mIL-1Ra3, or the IL-18R binding activity of a native sequence hIL-1Ra1, hIL-1Ra1L, or hIL-1Ra1V, which DNA has at least at or about 80% sequence identity, or at least at or about 85% sequence identity, or at least at or about 90% sequence identity, or at least at or about 95% sequence identity to (a) a DNA encoding an IL-11p, such as a mature IL-11p polypeptide, encoded by the cDNA insert in the vector deposited as ATCC Deposit No. 203588 (DNA85066-2534), ATCC Deposit No. 203587 (DNA96786-2534), ATCC Deposit No. 203589 (DNA96787-2534), ATCC Deposit No. 203590 (DNA92505-2534), ATCC Deposit No. 203846 (DNA102043-2534), or ATCC Deposit No. 203973 (DNA114876-2534), or (b) the complement of the DNA molecule of (a). In a preferred embodiment, the nucleic acid comprises a DNA encoding an IL-11p polypeptide, such as a mature IL-11p polypeptide, encoded by the cDNA insert in the vector deposited as ATCC Deposit No. 203588 (DNA85066-2534), ATCC Deposit No. 203587 (DNA96786-2534), ATCC Deposit No. 203586 (DNA92929-2534), ATCC Deposit No. 203589 (DNA96787-2534), ATCC Deposit No. 203590 (DNA92505-2534), ATCC Deposit No. 203846 (DNA102043-2534), ATCC Deposit No. 203973 (DNA114876-2534), or ATCC Deposit No. 203855 (DNA102044-2534).

In a further aspect, the invention concerns an isolated nucleic acid molecule comprising DNA encoding an IL-11p polypeptide that retains at least one biologic activity of a native sequence IL-1lp, such as the IL-1R binding activity of a native sequence hIL-1Ra3 or mIL-1Ra3, or the IL-18R binding activity of a native sequence hIL-1Ra1, hIL-1Ra1L, or hIL-1Ra1V, which DNA has at least at or about 80% sequence identity, or at least at or about 85% sequence identity, or at least at or about 90% sequence identity, or at least at or about 95% sequence identity to (a) a DNA encoding the entire amino acid sequence encoded by the longest open reading frame in the cDNA insert of a vector selected from the group consisting of the vectors deposited as ATCC Deposit No. 203588 (DNA85066-2534), ATCC Deposit No. 203587 (DNA96786-2534), ATCC Deposit No. 203589 (DNA96787-2534), ATCC Deposit No. 203590 (DNA92505-2534), ATCC Deposit No. 203846 (DNA102043-2534), and ATCC Deposit No. 203973 (DNA114876-2534), or (b) the complement of the DNA molecule of (a). In a preferred embodiment, the nucleic acid comprises (a) DNA encoding the entire amino acid sequence encoded by the longest open reading frame in the cDNA insert of a vector selected from the group consisting of the vectors deposited as ATCC Deposit No. 203588 (DNA85066-2534), ATCC Deposit No. 203587 (DNA96786-2534), ATCC Deposit No. 203586 (DNA92929-2534), ATCC Deposit No. 203589 (DNA96787-2534), and ATCC Deposit No. 203590 (DNA92505-2534), or (b) the complement of the DNA of (a). In another preferred embodiment, the nucleic acid comprises (a) DNA encoding the entire amino acid sequence encoded by the longest open reading frame in the cDNA insert of a vector selected from the group consisting of the vectors deposited as ATCC Deposit No. 203846 (DNA102043-2534), ATCC Deposit No. 203855 (DNA102044-2534), and ATCC Deposit No. 203973 (DNA114876-2534), or (b) the complement of the DNA of (a).

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In another aspect, the invention concerns an isolated nucleic acid molecule comprising DNA encoding an IL-11p polypeptide that retains at least one biologic activity of a native sequence IL-11p, such as the IL-1R binding activity of a native sequence IL-1Ra3 or mIL-1Ra3, or the IL-18R binding activity of a native sequence hIL-1Ra1, hIL-1Ra1L, or hIL-1Ra1V, which DNA has at least at or about 80% sequence identity, or at least at or about 85% sequence identity, or at least at or about 90% sequence identity, or at least at or about 95% sequence identity to (a) DNA encoding an amino acid sequence selected from the group consisting of: (1) the entire amino acid sequence encoded by the longest open reading frame in the cDNA insert in the vector deposited as ATCC Deposit No. 203588, (2) the entire amino acid sequence, or the entire amino acid sequence excluding the 36 N-terminal amino acid residues of such sequence, encoded by the longest open reading frame in the cDNA insert in the vector deposited as ATCC Deposit No. 203587, (3) the entire amino acid sequence, or the entire amino acid sequence excluding the Nterminal amino acid residue of such sequence, encoded by the longest open reading frame in the cDNA insert in the vector deposited as ATCC Deposit No. 203589, (4) the entire amino acid sequence, or the entire amino acid sequence excluding the N-terminal amino acid residue of such sequence, encoded by the longest open reading frame in the cDNA insert in the vector deposited as ATCC Deposit No. 203590, (5) the entire amino acid sequence, or the entire amino acid sequence excluding the N-terminal amino acid residue of such sequence, or the entire amino acid sequence excluding the 34 N-terminal amino acid residues of such sequence, encoded by the longest open reading frame in the cDNA insert in the vector deposited as ATCC Deposit No. 203846, and (6) the entire amino acid sequence, or the entire amino acid sequence excluding the N-terminal amino acid residue of such sequence, or the entire amino acid sequence excluding the 45 N-terminal amino acid residues of such sequence, encoded by the longest open reading frame in the cDNA insert in the vector deposited as ATCC Deposit No. 203973, or (b) the complement of the DNA of (a).

In a preferred embodiment, the nucleic acid comprises (a) DNA encoding an amino acid sequence selected from the group consisting of: (1) the entire amino acid sequence encoded by the longest open reading frame in the cDNA insert in the vector deposited as ATCC Deposit No. 203588, (2) the entire amino acid sequence, or the entire amino acid sequence excluding the 36 N-terminal amino acid residues of such sequence, encoded by the longest open reading frame in the cDNA insert in the vector deposited as ATCC Deposit No. 203587, (3) the entire amino acid sequence, or the entire amino acid sequence excluding the N-terminal amino acid residue of such sequence, encoded by the longest open reading frame in the cDNA insert in the vector deposited as ATCC Deposit No. 203589, (4) the entire amino acid sequence, or the entire amino acid sequence excluding the N-terminal amino acid residue of such sequence, encoded by the longest open reading frame in the cDNA insert in the vector deposited as ATCC Deposit No. 203590, (5) the entire amino acid sequence, or the entire amino acid sequence excluding the N-terminal amino acid residue of such sequence, or the entire amino acid sequence excluding the 34 Nterminal amino acid residues of such sequence, encoded by the longest open reading frame in the cDNA insert in the vector deposited as ATCC Deposit No. 203846, and (6) the entire amino acid sequence, or the entire amino acid sequence excluding the N-terminal amino acid residue

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of such sequence, or the entire amino acid sequence excluding the 45 N-terminal amino acid residues of such sequence, encoded by the longest open reading frame in the cDNA insert in the vector deposited as ATCC Deposit No. 203973, or (b) the complement of the DNA of (a).

In a still further aspect, the invention concerns an isolated nucleic acid molecule comprising (a) DNA encoding an IL-1lp polypeptide that retains at least one biologic activity of a native sequence IL-1lp, such as the IL-1RR binding activity of a native sequence hIL-1Ra1, or the IL-1R binding activity of a native sequence hIL-1Ra3, and which IL-1lp polypeptide has at least at or about 80% sequence identity, or at least at or about 85% sequence identity, or at least at or about 85% sequence identity, or at least at or about 95% sequence identity to an amino acid sequence selected from the group consisting of: (1) amino acid residues from at or about 37 to at or about 203, inclusive of Figure 2 (SEQ ID NO:5), (2) amino acid residues from at or about 34 to at or about 155, inclusive of Figure 7 (SEQ ID NO:13), (4) amino acid residues from at or about 34 to at or about 155, inclusive of Figure 9 (SEQ ID NO:16), (5) amino acid residues from at or about 26 to at or about 207, inclusive of Figure 15 (SEQ ID NO:16), (5) amino acid residues from at or about 26 to at or about 218, inclusive of Figure 19 (SEQ ID NO:25), or (b) the complement of the DNA of (a).

In a still further aspect, the invention concerns an isolated nucleic acid molecule comprising (a) DNA encoding an IL-11p polypeptide that retains at least one biologic activity of a native sequence IL-11p, such as the IL-18R binding activity of a native sequence hIL-1Ra1, or the IL-1R binding activity of a native sequence hIL-1Ra3 or mIL-1Ra3, and which IL-11p polypeptide has at least at or about 80% sequence identity, or at least at or about 85% sequence identity, or at least at or about 95% sequence identity to an amino acid sequence selected from the group consisting of: (1) amino acid residues from at or about 95 to at or about 134, inclusive of Figure 7 (SEQ ID NO:13), and (2) amino acid residues from at or about 95 to at or about 134, inclusive of Figure 9 (SEQ ID NO:16), or (b) the complement of the DNA of (a).

In a still further aspect, the invention concerns an isolated nucleic acid molecule comprising (a) DNA encoding an IL-1lp polypeptide that retains at least one biologic activity of a native sequence IL-1lp, such as the IL-18R binding activity of a native sequence hIL-1Ra1, or the IL-1R binding activity of a native sequence hIL-1Ra3 or mIL-1Ra3, and which IL-1lp polypeptide has at least at or about 80% sequence identity, or at least at or about 85% sequence identity, or at least at or about 95% sequence identity to the amino acid sequence of amino acid residues from at or about 80 to at or about 155, inclusive of Figure 7 (SEQ ID NO:13), or (b) the complement of the DNA of (a).

In a still further aspect, the invention concerns an isolated nucleic acid molecule comprising (a) DNA encoding an IL-11p polypeptide that retains at least one biologic activity of a native sequence IL-11p, such as the IL-18R binding activity of a native sequence hIL-1Ra1, or the IL-1R binding activity of a native sequence hIL-1Ra3 or mIL-1Ra3, and which IL-11p polypeptide has at least at or about 80% sequence identity, or at least at or about 85% sequence identity, or at least at or about 95% sequence

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identity to an amino acid sequence selected from the group consisting of: (1) amino acid residues from at or about 2 to at or about 155, inclusive of Figure 7 (SEQ ID NO:13), and (2) amino acid residues from at or about 2 to at or about 155, inclusive of Figure 9 (SEQ ID NO:16), or (b) the complement of the DNA of (a).

In a still further aspect, the invention concerns an isolated nucleic acid molecule comprising (a) DNA encoding an IL-11p polypeptide having at least at or about 80% sequence identity, or at least at or about 95% sequence identity, or at least at or about 95% sequence identity to an amino acid sequence selected from the group consisting of: (1) amino acid residues from at or about 95 to at or about 134, inclusive of Figure 7 (SEQ ID NO:13), and (2) amino acid residues from at or about 95 to at or about 134, inclusive of Figure 9 (SEQ ID NO:16), or (b) the complement of the DNA of (a).

In a still further aspect, the invention concerns an isolated nucleic acid molecule comprising (a) DNA encoding an IL-11p polypeptide having at least at or about 80% sequence identity, or at least at or about 85% sequence identity, or at least at or about 95% sequence identity to the amino acid sequence of amino acid residues from at or about 80 to at or about 155, inclusive of Figure 7 (SEQ ID NO:13), or (b) the complement of the DNA of (a).

In a still further aspect, the invention concerns an isolated nucleic acid molecule comprising (a) DNA encoding an IL-1lp polypeptide having at least at or about 80% sequence identity, or at least at or about 90% sequence identity, or at least at or about 90% sequence identity, or at least at or about 95% sequence identity to an amino acid sequence selected from the group consisting of: (1) amino acid residues from at or about 2 to at or about 155, inclusive of Figure 7 (SEQ ID NO:13), and (2) amino acid residues from at or about 2 to at or about 155, inclusive of Figure 9 (SEQ ID NO:16), or (b) the complement of the DNA of (a).

In a still further aspect, the invention concerns an isolated nucleic acid molecule comprising (a) DNA encoding an IL-11p polypeptide that retains at least one biologic activity of a native sequence IL-11p, such as the IL-18R binding activity of a native sequence hIL-1Ra1, hIL-1Ra1L, or hIL-1Ra1V, and which IL-11p polypeptide has at least at or about 80% sequence identity, or at least at or about 85% sequence identity, or at least at or about 90% sequence identity, or at least at or about 95% sequence identity to an amino acid sequence selected from the group consisting of: (1) amino acid residues from at or about 1 to at or about 207, inclusive of Figure 15 (SEQ 1D NO:19), and (2) amino acid residues from at or about 1 to at or about 218, inclusive of Figure 19 (SEQ ID NO:25), or (b) the complement of the DNA of (a).

In a further aspect, the invention concerns an isolated nucleic acid molecule comprising DNA encoding an IL-1lp that retains at least one biologic activity of a native sequence IL-1lp, such as the IL-1R binding activity of a native sequence hIL-1Ra3 or mIL-1Ra3, or the IL-1Ra binding activity of a native sequence hIL-1Ra1, hIL-1Ra1L, hIL-1Ra1S, or hIL-1Ra1V, which DNA is produced by hybridizing a test DNA molecule under stringent conditions with (a) a DNA molecule encoding an IL-1lp polypeptide selected from the group consisting of: (1) an IL-1lp polypeptide comprising the sequence of amino acid residues from at or about 37 to at or about 203, inclusive of Figure 2 (SEQ ID NO:5), (2) an IL-1lp polypeptide comprising the sequence of

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amino acid residues from at or about 15 to at or about 193, inclusive of Figure 3 (SEQ ID NO:7), (3) an IL-11p polypeptide comprising the sequence of amino acid residues from at or about 34 to at or about 155, inclusive of Figure 7 (SEQ ID NO:13), (4) an IL-11p polypeptide comprising the sequence of amino acid residues from at or about 34 to at or about 155, inclusive of Figure 9 (SEQ ID NO:16), (5) an IL-11p polypeptide comprising the sequence of amino acid residues from at or about 26 to at or about 207, inclusive of Figure 15 (SEQ ID NO:19), and (6) an IL-1lp polypeptide comprising the sequence of amino acid residues from at or about 46 to at or about 218, inclusive of Figure 19 (SEQ ID NO:25), or (b) the complement of the DNA molecule of (a), and, if the test DNA molecule encodes an IL-11p that retains at least one biologic activity of a native sequence IL-1lp, such as the IL-1R binding activity of a native sequence hIL-1Ra3 or mIL-1Ra3, or the IL-18R binding activity of a native sequence hIL-1Ra1, hIL-1Ra1L, or hIL-1Ra1V, and if the test DNA molecule has at least at or about an 80% sequence identity, or at least at or about an 85% sequence identity, or at least at or about a 90% sequence identity, or at least at or about a 95% sequence identity to the DNA molecule of (a) or (b), isolating the test DNA molecule.

In another aspect, the invention provides an isolated nucleic acid molecule comprising (a) DNA encoding an a polypeptide, such as IL-11p polypeptide, selected from the group consisting of: (1) a polypeptide, such as an hIL-1Ra1 polypeptide, comprising amino acid residues from at or about 37 to at or about 63, inclusive of Figure 2 (SEQ ID NO:5); (2) a polypeptide, such as an hIL-1Ra1 polypeptide, comprising amino acid residues from at or about 15 to at or about 53, inclusive of Figure 3 (SEQ ID NO:7); (3) a polypeptide, such as an hIL-1Ra2 polypeptide, comprising amino acid residues from at or about 1 to at or about 134, inclusive of Figure 5 (SEQ ID NO:10); (4) a polypeptide comprising amino acid residues from at or about 10 to at or about 134, inclusive of Figure 5 (SEQ ID NO:10); (5) a polypeptide, such as an hIL-1Ra2 polypeptide, consisting of amino acid residues from at or about 27 to at or about 134, inclusive of Figure 5 (SEQ ID NO:10); (6) a polypeptide, such as an hIL-1Ra2 fusion variant polypeptide, consisting of a native amino acid sequence of hIL-1Ra2 consisting of amino acid residues from at or about 27 to at or about 134, inclusive of Figure 5 (SEQ ID NO:10) fused at its N-terminus or C-terminus to a heterologous amino acid or amino acid sequence; (7) a polypeptide, such as an hIL-1Ra3 polypeptide, comprising amino acid residues from at or about 95 to at or about 134, inclusive of Figure 7 (SEO ID NO:13); and (8) a polypeptide, such as a mIL-1Ra3 polypeptide, comprising amino acid residues from at or about 95 to at or about 134, inclusive of Figure 9 (SEQ ID NO:16); or (b) the complement of the DNA of (a).

In another aspect, the invention provides an isolated nucleic acid molecule comprising (a) DNA encoding a polypeptide, such as an IL-1lp polypeptide, selected from the group consisting of: (1) a polypeptide, such as an hIL-1Ra1L polypeptide, comprising amino acid residues from at or about 26 to at or about 44, inclusive of Figure 15 (SEO ID NO:19); (2) a polypeptide, such as an hIL-1Ra1L polypeptide, comprising amino acid residues from at or about 1 to at or about 44, inclusive of Figure 15 (SEQ ID NO:19); (3) a polypeptide, such as an hIL-1Ra1L polypeptide, comprising amino acid residues from at or about 26 to at or about 78, inclusive of Figure 15 (SEQ ID NO:19); (4) a polypeptide, such as an hIL-1Ra1L polypeptide,

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comprising amino acid residues from at or about 1 to at or about 78, inclusive of Figure 15 (SEQ ID NO:19); (5) a polypeptide, such as an hIL-1Ra1S polypeptide, comprising amino acid residues from at or about 1 to at or about 38, inclusive of Figure 16 (SEQ ID NO:21); (6) a polypeptide, such as an hIL-1Ra1V polypeptide, comprising amino acid residues from at or about 37 to at or about 55, inclusive of Figure 19 (SEQ ID NO:25); (7) a polypeptide, such as an hIL-1Ra1V polypeptide, comprising amino acid residues from at or about 12 to at or about 55, inclusive of Figure 19 (SEQ ID NO:25); (8) a polypeptide, such as an hIL-1Ra1V polypeptide, comprising amino acid residues from at or about 1 to at or about 55, inclusive of Figure 19 (SEQ ID NO:25); (9) a polypeptide, such as an hIL-1Ra1V polypeptide, comprising amino acid residues from at or about 46 to at or about 55, inclusive of Figure 19 (SEQ ID NO:25); (10) a polypeptide, such as an hIL-1Ra1V polypeptide, comprising amino acid residues from at or about 46 to at or about 89, inclusive of Figure 19 (SEQ ID NO:25); (11) a polypeptide, such as an hIL-1Ra1V polypeptide, comprising amino acid residues from at or about 37 to at or about 89, inclusive of Figure 19 (SEQ ID NO:25); (12) a polypeptide, such as an hIL-1Ra1V polypeptide, comprising amino acid residues from at or about 12 to at or about 89, inclusive of Figure 19 (SEQ ID NO:25); and (13) a polypeptide, such as an hIL-1Ra1V polypeptide, comprising amino acid residues from at or about 1 to at or about 89, inclusive of Figure 19 (SEQ ID NO:25); or (b) the complement of the DNA of (a).

In another aspect, the invention provides an isolated nucleic acid molecule comprising (a) DNA encoding a polypeptide, such as an hIL-11p polypeptide, selected from the group consisting of: (1) a polypeptide, such as an hIL-1Ra1L polypeptide, consisting of a native amino acid sequence of hIL-1Ra1L consisting of amino acid residues from at or about 26 to at or about 207, inclusive of Figure 15 (SEQ ID NO:19) fused at its N-terminus or C-terminus to a heterologous amino acid or amino acid sequence; (2) a polypeptide, such as an hIL-1Ra1L polypeptide, consisting of a native amino acid sequence of hIL-1Ra1L consisting of amino acid residues from at or about 1 to at or about 207, inclusive of Figure 15 (SEQ ID NO:19) fused at its N-terminus or C-terminus to a heterologous amino acid or amino acid sequence; (3) a polypeptide, such as an hIL-1Ra1L polypeptide, consisting of amino acid residues from at or about 26 to at or about 207, inclusive of Figure 15 (SEQ ID NO:19); (4) a polypeptide, such as an hIL-1Ra1L polypeptide, consisting of amino acid residues from at or about 1 to at or about 207, inclusive of Figure 15 (SEQ ID NO:19); (5) a polypeptide, such as an hIL-1Ra1S fusion variant polypeptide, consisting of a native amino acid sequence of hIL-1Ra1S consisting of amino acid residues from at or about 26 to at or about 167, inclusive of Figure 16 (SEQ ID NO:21) fused at its N-terminus or C-terminus to a heterologous amino acid or amino acid sequence; (6) a polypeptide, such as an hIL-1Ra1S polypeptide, consisting of amino acid residues from at or about 26 to at or about 167, inclusive of Figure 16 (SEQ ID NO:21); (7) a polypeptide, such as an hIL-1Ra1S fusion variant polypeptide, consisting of a native amino acid sequence of hIL-1Ra1S consisting of amino acid residues from at or about 1 to at or about 167, inclusive of Figure 16 (SEQ ID NO:21) fused at its N-terminus or C-terminus to a heterologous amino acid or amino acid sequence; (8) a polypeptide, such as an hIL-1Ra1S polypeptide, consisting of amino acid residues from at or about 1 to at or about 167, inclusive of Figure 16 (SEQ ID NO:21);

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(9) a polypeptide, such as an hIL-1Ra1S fusion variant polypeptide, consisting of a native amino acid sequence of hIL-1Ra1S consisting of amino acid residues from at or about 39 to at or about 167, inclusive of Figure 16 (SEQ ID NO:21) fused at its N-terminus or C-terminus to a heterologous amino acid or amino acid sequence; (10) a polypeptide, such as an hIL-1Ra1S fusion variant polypeptide, consisting of a native amino acid sequence of hIL-1Ra1S consisting of amino acid residues from at or about 47 to at or about 167, inclusive of Figure 16 (SEQ ID NO:21) fused at its N-terminus or C-terminus to a heterologous amino acid or amino acid sequence; (11) a polypeptide, such as an hIL-1Ra1S polypeptide, consisting of amino acid residues from at or about 39 to at or about 167, inclusive of Figure 16 (SEQ ID NO:21); (12) a polypeptide, such as an hIL-1Ra1S polypeptide, consisting of amino acid residues from at or about 47 to at or about 167, inclusive of Figure 16 (SEQ ID NO:21); (13) a polypeptide, such as an hIL-1Ra1V polypeptide, consisting of a native amino acid sequence of hIL-1Ra1V consisting of amino acid residues from at or about 1 to at or about 218, inclusive of Figure 19 (SEQ ID NO:25) fused at its N-terminus or C-terminus to a heterologous amino acid or amino acid sequence; (14) a polypeptide, such as an hIL-1Ra1V polypeptide, consisting of amino acid residues from at or about 1 to at or about 218, inclusive of Figure 19 (SEQ ID NO:25); (15) a polypeptide, such as an hIL-1Ra1V polypeptide, consisting of a native amino acid sequence of hIL-1Ra1V consisting of amino acid residues from at or about 12 to at or about 218, inclusive of Figure 19 (SEQ ID NO:25) fused at its N-terminus or C-terminus to a heterologous amino acid or amino acid sequence; (16) a polypeptide, such as an hIL-1Ra1V polypeptide, consisting of amino acid residues from at or about 12 to at or about 218, inclusive of Figure 19 (SEQ ID NO:25); (17) a polypeptide, such as an hIL-1Ra1V fusion variant polypeptide, consisting of a native amino acid sequence of hIL-1Ra1V consisting of amino acid residues from at or about 37 to at or about 218, inclusive of Figure 19 (SEQ ID NO:25) fused at its N-terminus or C-terminus to a heterologous amino acid or amino acid sequence; (18) a polypeptide, such as an hIL-1Ra1V fusion variant polypeptide, consisting of a native amino acid sequence of hIL-1Ra1V consisting of amino acid residues from at or about 46 to at or about 218, inclusive of Figure 19 (SEQ ID NO:25) fused at its N-terminus or C-terminus to a heterologous amino acid or amino acid sequence; (19) a polypeptide, such as an hIL-1Ra1V polypeptide, consisting of amino acid residues from at or about 37 to at or about 218, inclusive of Figure 19 (SEQ ID NO:25); and (20) a polypeptide, such as an hIL-1Ra1V polypeptide, consisting of amino acid residues from at or about 46 to at or about 218, inclusive of Figure 19 (SEQ ID NO:25); or (b) the complement of the DNA of (a).

In another aspect, the invention provides an isolated DNA molecule selected from the group consisting of: (1) a DNA molecule encoding a polypeptide, such as an hIL-1Ra1 polypeptide, comprising the amino acid sequence of amino acid residues from at or about 37 to at or about 203, inclusive of Figure 2 (SEQ ID NO:5); (2) a DNA molecule encoding a polypeptide, such as an hIL-1Ra1 polypeptide, comprising the amino acid sequence of amino acid residues from at or about 15 to at or about 193, inclusive of Figure 3 (SEQ ID NO:7); (3) a DNA molecule encoding a polypeptide, such as an hIL-1Ra2 polypeptide, comprising the amino acid sequence of amino acid residues from at or about 1 to at or about 134, inclusive of Figure 5 (SEQ ID

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NO:10); (4) a DNA molecule encoding a polypeptide comprising the amino acid sequence of amino acid residues from at or about 10 to at or about 134, inclusive of Figure 5 (SEQ ID NO:10); (5) a DNA molecule encoding a polypeptide, such as an hIL-1Ra2 fusion variant polypeptide, consisting of a native amino acid sequence of hIL-1Ra2 consisting of amino acid residues from at or about 27 to at or about 134, inclusive of Figure 5 (SEQ ID NO:10) fused at its N-terminus or C-terminus to a heterologous amino acid or amino acid sequence; (6) a DNA molecule encoding a polypeptide, such as an hIL-1Ra2 polypeptide, consisting of the amino acid sequence of amino acid residues from at or about 27 to at or about 134, inclusive of Figure 5 (SEQ ID NO:10); (7) a DNA molecule encoding a polypeptide, such as an hIL-1Ra3 polypeptide, comprising the amino acid sequence of amino acid residues from at or about 95 to at or about 134, inclusive of Figure 7 (SEQ ID NO:13); (8) a DNA molecule encoding a polypeptide, such as a mIL-1Ra3 polypeptide, comprising the amino acid sequence of amino acid residues from at or about 134, inclusive of Figure 9 (SEQ ID NO:16); and (9) the complement of any of the DNA molecules of (1)-(8).

In another aspect, the invention provides an isolated DNA molecule selected from the group consisting of: (1) a DNA molecule encoding a polypeptide, such as an hIL-1Ra1L polypeptide, comprising the amino acid sequence of amino acid residues from at or about 26 to at or about 207, inclusive of Figure 15 (SEQ ID NO:19); (2) a DNA molecule encoding a polypeptide, such as an hIL-1Ra1S polypeptide, comprising the amino acid sequence of amino acid residues from at or about 26 to at or about 167, inclusive of Figure 16 (SEQ ID NO:21); (3) a DNA molecule encoding a polypeptide, such as an hIL-1Ra1V polypeptide, comprising the amino acid sequence of amino acid residues from at or about 37 to at or about 218, inclusive of Figure 19 (SEQ ID NO:25); (4) a DNA molecule encoding a polypeptide, such as an hIL-1Ra1V polypeptide, comprising the amino acid sequence of amino acid residues from at or about 46 to at or about 218, inclusive of Figure 19 (SEQ ID NO:25); and (5) the complement of any of the DNA molecules of (1)-(4).

In another aspect, the invention provides an isolated DNA molecule selected from the group consisting of: (1) a DNA molecule encoding a polypeptide, such as an hIL-1Ra1L polypeptide, comprising the amino acid sequence of amino acid residues from at or about 1 to at or about 207, inclusive of Figure 15 (SEQ ID NO:19); (2) a DNA molecule encoding a polypeptide, such as an hIL-1Ra1S polypeptide, comprising the amino acid sequence of amino acid residues from at or about 1 to at or about 167, inclusive of Figure 16 (SEQ ID NO:21); (3) a DNA molecule encoding a polypeptide, such as an hIL-1Ra1V polypeptide, comprising the amino acid, sequence of amino acid residues from at or about 218, inclusive of Figure 19 (SEQ ID NO:25); (4) a DNA molecule encoding a polypeptide, such as an hIL-1Ra1V polypeptide, comprising the amino acid sequence of amino acid residues from at or about 1 to at or about 218, inclusive of Figure 19 (SEQ ID NO:25); and (5) the complement of any of the DNA molecules of (1)-(4).

In another aspect, the invention provides an isolated DNA molecule selected from the group consisting of: (1) a DNA molecule encoding a polypeptide, such as an hIL-1Ra3

WO 00/39297 PCT/US99/30720

polypeptide, comprising amino acid residues from at or about 80 to at or about 155, inclusive of Figure 7 (SEQ ID NO:13); and (2) the complement of the DNA molecule of (1).

In another aspect, the invention provides an isolated DNA molecule selected from the group consisting of: (1) a DNA molecule encoding a polypeptide, such as an hIL-1Ra1 polypeptide, comprising the amino acid sequence of amino acid residues from at or about 1 to at or about 203, inclusive of Figure 2 (SEQ ID NO:5); (2) a DNA molecule encoding a polypeptide, such as an hIL-1Ra1 polypeptide, comprising the amino acid sequence of amino acid residues from at or about 1 to at or about 193, inclusive of Figure 3 (SEQ ID NO:7); (3) a DNA molecule encoding a polypeptide, such as an hIL-1Ra3 polypeptide, comprising the amino acid sequence of amino acid residues from at or about 34 to at or about 155, inclusive of Figure 7 (SEQ ID NO:13); (4) a DNA molecule encoding a polypeptide, such as a mIL-1Ra3 polypeptide, comprising the amino acid sequence of amino acid residues from at or about 34 to at or about 155, inclusive of Figure 9 (SEQ ID NO:16); and (5) the complement of any of the DNA molecules of (1)-(4).

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In another aspect, the invention provides an isolated DNA molecule selected from the group consisting of: (1) a DNA molecule encoding a polypeptide, such as an hIL-1Ra3 polypeptide, comprising the amino acid sequence of amino acid residues from at or about 1 to at or about 155, inclusive of Figure 7 (SEQ ID NO:13); (2) a DNA molecule encoding a polypeptide, such as a mIL-1Ra3 polypeptide, comprising the amino acid sequence of amino acid residues from at or about 1 to at or about 155, inclusive of Figure 9 (SEQ ID NO:16); and (3) the complement of any of the DNA molecules of (1)-(2).

In another aspect, the invention provides an isolated DNA molecule selected from the group consisting of: (1) a DNA molecule encoding a polypeptide, such as an hIL-1Ra3 polypeptide, comprising the amino acid sequence of amino acid residues from at or about 2 to at or about 155, inclusive of Figure 7 (SEQ ID NO:13); (2) a DNA molecule encoding a polypeptide, such as a mIL-1Ra3 polypeptide, comprising the amino acid sequence of amino acid residues from at or about 2 to at or about 155, inclusive of Figure 9 (SEQ ID NO:16); and (3) the complement of any of the DNA molecules of (1)-(2).

In another aspect, the invention provides an isolated nucleic acid molecule comprising (a) a DNA molecule encoding a polypeptide selected from the group consisting of: (1) a polypeptide comprising an hIL-IRa1 polypeptide, such as a mature hIL-IRa1 polypeptide, encoded by the cDNA insert in the vector deposited as ATCC Deposit No. 203588; (2) a polypeptide comprising an hIL-IRa1 polypeptide, such as a mature hIL-IRa1 polypeptide, encoded by the cDNA insert in the vector deposited as ATCC Deposit No. 203587; (3) a polypeptide consisting of an hIL-IRa2 polypeptide, such as a mature hIL-IRa2 polypeptide, encoded by the cDNA insert in the vector deposited as ATCC Deposit No. 203586, which hIL-IRa2 polypeptide is fused at its N-terminus or C-terminus to a heterologous amino acid or amino acid sequence; (4) a polypeptide consisting of an hIL-IRa2 polypeptide, such as a mature hIL-IRa2 polypeptide, encoded by the cDNA insert in the vector deposited as ATCC Deposit No. 203586; (6) a polypeptide comprising an hIL-IRa3 polypeptide, such as a mature mIL-IRa3 polypeptide, encoded by the cDNA insert in the vector deposited as ATCC Deposit No. 203589; and (6) a polypeptide comprising a mIL-IRa3 polypeptide, such as a mature mIL-IRa3

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polypeptide, encoded by the cDNA insert in the vector deposited as ATCC Deposit No. 203590; or (b) the complement of the DNA molecule of (a).

In another aspect, the invention provides an isolated nucleic acid molecule comprising (a) a DNA molecule encoding a polypeptide selected from the group consisting of: (1) a polypeptide comprising an hIL-1Ra1L polypeptide, such as a mature hIL-1Ra1L polypeptide, encoded by the cDNA insert in the vector deposited as ATCC Deposit No.203846; (2) a polypeptide consisting of an hIL-1Ra1S polypeptide, such as a mature hIL-1Ra1S polypeptide, encoded by the cDNA insert in the vector deposited as ATCC Deposit No.203855, which hIL-1Ra1S polypeptide is fused at its N-terminus or C-terminus to a heterologous amino acid or amino acid sequence; (3) a polypeptide consisting of an hIL-1Ra1S polypeptide, such as a mature hIL-1Ra1S polypeptide, encoded by the cDNA insert in the vector deposited as ATCC Deposit No.203855; (4) a polypeptide comprising an hIL-1Ra1V polypeptide, encoded by the cDNA insert in the vector deposited as ATCC Deposit No.203973; or (b) the complement of the DNA molecule of (a).

In another aspect, the invention provides an isolated nucleic acid molecule comprising (a) a DNA molecule encoding a polypeptide comprising an hIL-1Ra1S polypeptide, such as a mature hIL-1Ra1S polypeptide, encoded by the cDNA insert in the vector deposited as ATCC Deposit No. 203855; or (b) the complement of the DNA molecule of (a).

In another aspect, the invention provides an isolated nucleic acid molecule comprising (a) a DNA encoding a polypeptide selected from the group consisting of: (1) a polypeptide comprising the entire amino acid sequence encoded by the longest open reading frame in the cDNA insert in the vector deposited as ATCC Deposit No. 203588; (2) a polypeptide comprising the entire amino acid sequence, or the entire amino acid sequence excluding the 36 N-terminal amino acid residues of such sequence, encoded by the longest open reading frame in the cDNA insert in the vector deposited as ATCC Deposit No. 203587; (3) a polypeptide comprising the entire amino acid sequence, or the entire amino acid sequence excluding the N-terminal amino acid residue of such sequence, or the entire amino acid sequence excluding the 9 N-terminal amino acid residues of such sequence, encoded by the longest open reading frame in the cDNA insert in the vector deposited as ATCC Deposit No. 203586; (4) a polypeptide comprising the entire amino acid sequence, or the entire amino acid sequence excluding the N-terminal amino acid residue of such sequence, encoded by the longest open reading frame in the cDNA insert in the vector deposited as ATCC Deposit No. 203589; and (5) a polypeptide comprising the entire amino acid sequence, or the entire amino acid sequence excluding the N-terminal amino acid residue of such sequence, encoded by the longest open reading frame in the cDNA insert in the vector deposited as ATCC Deposit No. 203590; or (b) the complement of the DNA of (a).

In another aspect, the invention provides an isolated nucleic acid molecule comprising (a) a DNA encoding a polypeptide comprising the entire amino acid sequence encoded by the longest open reading frame in the cDNA insert of a vector selected from the group consisting of the vectors deposited as ATCC Deposit Nos. 203588, 203586,203589, 203590, and 203973, or (b) the complement of the DNA of (a).

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In another aspect, the invention provides an isolated nucleic acid molecule comprising (a) a DNA encoding a polypeptide selected from the group consisting of: (1) a polypeptide comprising the entire amino acid sequence, or the entire amino acid sequence excluding the Nterminal amino acid residue of such sequence, or the entire amino acid sequence excluding the 34 N-terminal amino acid residues of such sequence, encoded by the longest open reading frame in the cDNA insert in the vector deposited as ATCC Deposit No. 203846; (2) a polypeptide comprising the entire amino acid sequence, or the entire amino acid sequence excluding the Nterminal amino acid residue of such sequence, or the entire amino acid sequence excluding the 25 N-terminal amino acid residues of such sequence, encoded by the longest open reading frame in the cDNA insert in the vector deposited as ATCC Deposit No. 204855; and (3) a polypeptide comprising the entire amino acid sequence, or the entire amino acid sequence excluding the Nterminal amino acid residue of such sequence, or the entire amino acid sequence excluding the 11 N-terminal amino acid residues of such sequence, or the entire amino acid sequence excluding the 36 N-terminal amino acid residues of such sequence, or the entire amino acid sequence excluding the 45 N-terminal amino acid residues of such sequence, encoded by the longest open reading frame in the cDNA insert in the vector deposited as ATCC Deposit No. 203973; or (b) the complement of the DNA of (a).

In another aspect, the invention provides an isolated nucleic acid molecule comprising (a) a DNA encoding a non-naturally occurring, chimeric polypeptide formed by fusing the entire amino acid sequence excluding the 38 N-terminal amino acid residues of such sequence, or the entire amino acid sequence excluding the 46 N-terminal amino acid residues of such sequence, encoded by the longest open reading frame in the cDNA insert in the vector deposited as ATCC Deposit No. 203855, at its N-terminus or C-terminus to a heterologous amino acid or amino acid sequence; or (b) the complement of the DNA of (a).

In another aspect, the invention provides an isolated nucleic acid molecule comprising (a) a DNA encoding a polypeptide consisting of the entire amino acid sequence excluding the 38 N-terminal amino acid residues of such sequence, or the entire amino acid sequence excluding the 46 N-terminal amino acid residues of such sequence, encoded by the longest open reading frame in the cDNA insert in the vector deposited as ATCC Deposit No. 203855; or (b) the complement of the DNA of (a).

In another aspect, the invention provides an isolated nucleic acid molecule comprising (a) a DNA molecule encoding a polypeptide comprising the entire amino acid sequence encoded by the longest open reading frame in the cDNA insert of a vector selected from the group consisting of the vectors deposited as ATCC Deposit Nos. 203846, 203855 and 203973, or (b) the complement of the DNA of (a).

In another aspect, the invention provides an isolated DNA molecule selected from the group consisting of: (1) a DNA molecule which encodes a polypeptide, such as an hIL-1Ra1V fusion variant polypeptide, consisting of a native amino acid sequence of hIL-1Ra1V consisting of amino acid residues from at or about 37 to at or about 218, inclusive of Figure 19 (SEQ ID NO:25) fused at its N-terminus or C-terminus to a heterologous amino acid or amino acid sequence, and which DNA molecule comprises the nucleic acid sequence in the sense strand of

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Figure 19 (SEQ ID NO:24) that encodes the native amino acid sequence; (2) a DNA molecule which encodes a polypeptide, such as an hIL-1Ra1V fusion variant polypeptide, consisting of a native amino acid sequence of hlL-1Ra1V consisting of amino acid residues from at or about 46 to at or about 218, inclusive of Figure 19 (SEQ ID NO:25) fused at its N-terminus or C-terminus to a heterologous amino acid or amino acid sequence, and which DNA molecule comprises the nucleic acid sequence in the sense strand of Figure 19 (SEO ID NO:24) that encodes the native amino acid sequence; (3) a DNA molecule which encodes a polypeptide, such as an hIL-1Ra1V polypeptide, consisting of a native amino acid sequence of hIL-1Ra1V consisting of amino acid residues from at or about 37 to at or about 218, inclusive of Figure 19 (SEQ ID NO:25), and which DNA molecule comprises the nucleic acid sequence in the sense strand of Figure 19 (SEQ ID NO:24) that encodes the native amino acid sequence; (4) a DNA molecule which encodes a polypeptide, such as an hlL-1Ra1V polypeptide, consisting of a native amino acid sequence of hIL-1Ra1V consisting of amino acid residues from at or about 46 to at or about 218, inclusive of Figure 19 (SEQ ID NO:25), and which DNA molecule comprises the nucleic acid sequence in the sense strand of Figure 19 (SEQ ID NO:24) that encodes the native amino acid sequence; and (5) the complement of any of the DNA molecules of (1)-(4).

In another aspect, the invention provides an isolated DNA molecule selected from the group consisting of: (1) a DNA molecule which encodes a polypeptide, such as an hlL-1Ra1 polypeptide, and which DNA molecule comprises the nucleic acid sequence of nucleotide positions from at or about 118 to at or about 618, inclusive in the sense strand of Figure 2 (SEQ ID NO:4); (2) a DNA molecule which encodes a polypeptide, such as an hIL-1Ra1 polypeptide, and which DNA molecule comprises the nucleic acid sequence of nucleotide positions from at or about 145 to at or about 681, inclusive in the sense strand of Figure 3 (SEQ ID NO:6); (3) a DNA molecule which encodes a polypeptide, such as an hIL-1Ra2 polypeptide, and which DNA molecule comprises the nucleic acid sequence of nucleotide positions from at or about 96 to at or about 497, inclusive in the sense strand of Figure 5 (SEQ ID NO:9); (4) a DNA molecule which comprises the nucleic acid sequence of nucleotide positions from at or about 123 to at or about 497, inclusive in the sense strand of Figure 5 (SEQ ID NO:9); (5) a DNA molecule which encodes a polypeptide, such as an hIL-1Ra2 fusion variant polypeptide, consisting of a native amino acid sequence of hIL-1Ra2 consisting of amino acid residues from at or about 27 to at or about 134, inclusive of Figure 5 (SEQ ID NO:10) fused at its N-terminus or C-terminus to a heterologous amino acid or amino acid sequence, and which DNA molecule comprises the nucleic acid sequence in the sense strand of Figure 5 (SEQ ID NO:9) that encodes the native amino acid sequence; (6) a DNA molecule which encodes a polypeptide, such as an hIL-1Ra2 polypeptide, consisting of a native amino acid sequence of hlL-1Ra2 consisting of amino acid residues from at or about 27 to at or about 134, inclusive of Figure 5 (SEQ ID NO:10), and which DNA molecule comprises the nucleic acid sequence in the sense strand of Figure 5 (SEQ ID NO:9) that encodes the native amino acid sequence; (7) a DNA molecule which encodes a polypeptide, such as an hIL-1Ra3 polypeptide, and which DNA molecule comprises the nucleic acid sequence of nucleotide positions from at or about 283 to at or about 402, inclusive in the sense strand of Figure 7 (SEQ ID NO:12); (8) a DNA molecule which encodes a polypeptide, such as a mIL-1Ra3

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polypeptide, and which DNA molecule comprises the nucleic acid sequence of nucleotide positions from at or about 427 to at or about 546, inclusive in the sense strand of Figure 9 (SEQ ID NO:15); and (9) the complement of any of the DNA molecules of (1)-(8).

In another aspect, the invention provides an isolated DNA molecule selected from the group consisting of: (1) a DNA molecule which encodes a polypeptide, such as an hIL-1Ra1L polypeptide, and which DNA molecule comprises the nucleic acid sequence of nucleotide positions from at or about 79 to at or about 624, inclusive in the sense strand of Figure 15 (SEQ ID NO:18]; (2) a DNA molecule which encodes a polypeptide, such as an hIL-1Ra1S polypeptide, and which DNA molecule comprises the nucleic acid sequence of nucleotide positions from at or about 79 to at or about 504, inclusive in the sense strand of Figure 16 (SEQ ID NO:20); (3) a DNA molecule which encodes a polypeptide, such as an hIL-1Ra1S fusion variant polypeptide, consisting of a native amino acid sequence of hIL-1Ra1S consisting of amino acid residues from at or about 39 to at or about 167, inclusive of Figure 16 (SEQ ID NO:21) fused at its N-terminus or C-terminus to a heterologous amino acid or amino acid sequence, and which DNA molecule comprises the nucleic acid sequence in the sense strand of Figure 16 (SEQ ID NO:20) that encodes the native amino acid sequence; (4) a DNA molecule which encodes a polypeptide, such as an hIL-1Ra1S fusion variant polypeptide, consisting of a native amino acid sequence of hIL-1Ra1S consisting of amino acid residues from at or about 47 to at or about 167, inclusive of Figure 16 (SEQ ID NO:21) fused at its N-terminus or C-terminus to a heterologous amino acid or amino acid sequence, and which DNA molecule comprises the nucleic acid sequence in the sense strand of Figure 16 (SEQ ID NO:20) that encodes the native amino acid sequence; (5) a DNA molecule which encodes a polypeptide, such as an hIL-1Ra1S polypeptide, consisting of a native amino acid sequence of hIL-1Ra1S consisting of amino acid residues from at or about 39 to at or about 167, inclusive of Figure 16 (SEQ ID NO:21), and which DNA molecule comprises the nucleic acid sequence in the sense strand of Figure 16 (SEQ ID NO:20) that encodes the native amino acid sequence; (6) a DNA molecule which encodes a polypeptide, such as an hlL-1Ra1S polypeptide, consisting of a native amino acid sequence of hIL-1Ra1S consisting of amino acid residues from at or about 47 to at or about 167, inclusive of Figure 16 (SEQ ID NO:21), and which DNA molecule comprises the nucleic acid sequence in the sense strand of Figure 16 (SEQ ID NO:20) that encodes the native amino acid sequence; (7) a DNA molecule which encodes a polypeptide, such as an hIL-1Ra1V polypeptide, and which DNA molecule comprises the nucleic acid sequence of nucleotide positions from at or about 181 to at or about 729, inclusive in the sense strand of Figure 19 (SEQ ID NO:24); (8) a DNA molecule which encodes a polypeptide, such as an hlL-1Ra1V polypeptide, and which DNA molecule comprises the nucleic acid sequence of nucleotide positions from at or about 208 to at or about 729, inclusive in the sense strand of Figure 19 (SEQ ID NO:24); and (9) the complement of any of the DNA molecules of (1)-(8).

In another aspect, the invention provides an isolated DNA molecule selected from the group consisting of: (1) a DNA molecule which encodes a polypeptide, such as an hIL-1Ra1L polypeptide, and which DNA molecule comprises the nucleic acid sequence of nucleotide positions from at or about 4 to at or about 624, inclusive in the sense strand of Figure 15 (SEQ ID NO:18); (2) a DNA molecule which encodes a polypeptide, such as an hIL-1Ra1S polypeptide,

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and which DNA molecule comprises the nucleic acid sequence of nucleotide positions from at or about 4 to at or about 504, inclusive in the sense strand of Figure 16 (SEQ ID NO:20); (3) a DNA molecule which encodes a polypeptide, such as an hIL-1Ra1V polypeptide, and which DNA molecule comprises the nucleic acid sequence of nucleotide positions from at or about 106 to at or about 729, inclusive in the sense strand of Figure 19 (SEO ID NO:24); (4) a DNA molecule which encodes a polypeptide, such as an hIL-1Ra1 V polypeptide, and which comprises the nucleic acid sequence of nucleotide positions from at or about 73 to at or about 729, inclusive in the sense strand of Figure 19 (SEQ ID NO:24); and (5) the complement of any of the DNA molecules of (1)-(4).

In another aspect, the invention provides an isolated DNA molecule selected from the group consisting of: (1) a DNA molecule comprising the nucleic acid sequence of nucleotide positions from at or about 103 to at or about 681, inclusive in the sense strand of Figure 3 (SEQ ID NO:6); (2) a DNA molecule comprising the nucleic acid sequence of nucleotide positions from at or about 100 to at or about 465, inclusive in the sense strand of Figure 7 (SEQ ID NO:12); (3) a DNA molecule comprising the nucleic acid sequence of nucleotide positions from at or about 244 to at or about 609, inclusive in the sense strand of Figure 9 (SEQ ID NO:15); and (4) the complement of any of the DNA molecules of (1)-(3).

In another aspect, the invention provides an isolated DNA molecule comprising (a) the complete DNA sequence in the sense strand of Figure 2 (SEQ ID NO:4), Figure 3 (SEQ ID NO:6), Figure 5 (SEQ ID NO:9), Figure 7 (SEQ ID NO:12), or Figure 9 (SEQ ID NO:15), or (b) the complement of (a).

In another aspect, the invention provides an isolated DNA molecule comprising (a) the complete DNA sequence in the sense strand of Figure 15 (SEO ID NO:18), Figure 16 (SEQ ID NO:20), or Figure 19 (SEO ID NO:24), or (b) the complement of (a).

In a specific aspect, the invention provides an isolated nucleic acid molecule comprising DNA encoding an IL-11p polypeptide, with or without the N-terminal signal sequence and/or the initiating methionine, or is complementary to such IL-1lp encoding nucleic acid molecule. The signal peptide has been tentatively identified as extending from amino acid position 1 to at or about amino acid position 14 in the IL-1lp sequence of Figure 3 (SEQ ID NO:7), from amino acid position 1 to at or about amino acid position 26 in the IL-1lp sequence of Figure 5 (SEQ ID NO:10), from amino acid position 1 to at or about amino acid position 33 in the IL-1lp sequence of Figure 7 (SEQ ID NO:13), and from amino acid position 1 to at or about amino acid position 33 in the IL-11p sequence of Figure 9 (SEQ ID NO:16).

The IL-1lp sequence of amino acids from at or about 1 to at or about 207 of Figure 15 (SEQ ID NO:19) is believed to behave as a mature sequence (without a presequence that is removed in post-translational processing) in certain animal cells. In addition, it is believed that other animal cells recognize and remove in post-translational processing one or more signal peptide(s) contained in the sequence of amino acid positions 1 to about 34 of Figure 15 (SEQ ID NO:19).

The IL-1lp sequence of amino acids from at or about 1 to at or about 167 of Figure 16 (SEQ ID NO:21) is believed to behave as a mature sequence (without a presequence that is

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removed in post-translational processing) in certain animal cells. In addition, it is believed that other animal cells recognize and remove in post-translational processing one or more signal peptide(s) contained in the sequence of amino acid positions 1 to about 46 in the IL-1lp sequence of Figure 16 (SEO ID NO:21).

The IL-1lp sequence of amino acids from at or about 1 to at or about 218 of Figure 19 (SEO ID NO:25) is believed to behave as a mature sequence (without a presequence that is removable in post-translational processing) in certain animal cells. The IL-11p sequence of amino acids from at or about 12 to at or about 218 of Figure 19 (SEQ ID NO:25) that results from initiation of translation at the start codon occurring at nucleotide positions 106-108 is also believed to behave as mature sequence in certain animal cells. It is further believed that other animal cells recognize and remove in post-translational processing one or more signal peptide(s) contained in the sequence of amino acid positions 1 to 45 in the IL-1lp polypeptide of amino acid positions 1 to 218 of Figure 19 (SEO ID NO:25) or contained in the sequence of amino acid positions 12 to 45 in the IL-1lp polypeptide of amino acid positions 12 to 218 of Figure 19 (SEQ ID NO:25).

In a still further aspect, the invention concerns an isolated nucleic acid molecule comprising (a) DNA encoding an IL-1lp polypeptide that retains at least one biologic activity of a native sequence IL-1lp, such as the IL-18R binding activity of a native sequence hIL-1Ra1, or the IL-1R binding activity of a native sequence hIL-1Ra3 or mIL-1Ra3, and which IL-1lp polypeptide has at least at or about 80% positives, or at least at or about 85% positives, or at least at or about 90% positives, or at least at or about 95% positives to an amino acid sequence selected from the group consisting of: (1) amino acid residues from at or about 37 to at or about 203, inclusive of Figure 2 (SEQ ID NO:5), (2) amino acid residues from at or about 15 to at or about 193, inclusive of Figure 3 (SEQ ID NO:7), (3) amino acid residues from at or about 34 to at or about 155, inclusive of Figure 7 (SEQ ID NO:13), (4) amino acid residues from at or about 34 to at or about 155, inclusive of Figure 9 (SEQ ID NO:16), (5) amino acid residues from at or about 26 to at or about 207, inclusive of Figure 15 (SEQ ID NO:19), and (6) amino acid residues from at or about 46 to at or about 218, inclusive of Figure 19 (SEQ ID NO:25), or (b) the complement of the DNA of (a).

In a still further aspect, the invention concerns an isolated nucleic acid molecule comprising (a) DNA encoding an IL-1lp polypeptide that retains at least one biologic activity of a native sequence IL-1lp, such as the IL-18R binding activity of a native sequence hIL-1Ra1, or the IL-1R binding activity of a native sequence hIL-1Ra3 or mIL-1Ra3, and which IL-1lp polypeptide has at least at or about 80% positives, or at least at or about 85% positives, or at least at or about 90% positives, or at least at or about 95% positives to an amino acid sequence selected from the group consisting of: (1) amino acid residues from at or about 95 to at or about 134, inclusive of Figure 7 (SEO ID NO:13), and (2) amino acid residues from at or about 95 to at or about 134, inclusive of Figure 9 (SEQ ID NO:16), or (b) the complement of the DNA of (a).

In a still further aspect, the invention concerns an isolated nucleic acid molecule comprising (a) DNA encoding an IL-1lp polypeptide that retains at least one biologic activity of a native sequence IL-1lp, such as the IL-18R binding activity of a native sequence hIL-1Ra1, or

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the IL-1R binding activity of a native sequence hIL-1Ra3 or mIL-1Ra3, and which IL-1lp polypeptide has at least at or about 80% positives, or at least at or about 85% positives, or at least at or about 90% positives, or at least at or about 95% positives to the amino acid sequence of amino acid residues from at or about 80 to at or about 155, inclusive of Figure 7 (SEQ ID NO:13), or (b) the complement of the DNA of (a).

In a still further aspect, the invention concerns an isolated nucleic acid molecule comprising (a) DNA encoding an IL-11p polypeptide that retains at least one biologic activity of a native sequence IL-11p, such as the IL-18R binding activity of a native sequence hIL-1Ra1, or the IL-1R binding activity of a native sequence hIL-1Ra3 or mIL-1Ra3, and which IL-11p polypeptide has at least at or about 80% positives, or at least at or about 85% positives, or at least at or about 95% positives to an amino acid sequence selected from the group consisting of: (1) amino acid residues from at or about 2 to at or about 155, inclusive of Figure 7 (SEQ ID NO:13), and (2) amino acid residues from at or about 2 to at or about 155, inclusive of Figure 9 (SEQ ID NO:16), or (b) the complement of the DNA of (a).

In a still further aspect, the invention concerns an isolated nucleic acid molecule comprising (a) DNA encoding an IL-11p polypeptide having at least at or about 80% positives, or at least at or about 85% positives, or at least at or about 90% positives, or at least at or about 95% positives to an amino acid sequence selected from the group consisting of: (1) amino acid residues from at or about 95 to at or about 134, inclusive of Figure 7 (SEQ ID NO:13), and (2) amino acid residues from at or about 95 to at or about 134, inclusive of Figure 9 (SEQ ID NO:16), or (b) the complement of the DNA of (a).

In a still further aspect, the invention concerns an isolated nucleic acid molecule comprising (a) DNA encoding an IL-11p polypeptide having at least at or about 80% positives, or at least at or about 85% positives, or at least at or about 90% positives, or at least at or about 95% positives to the amino acid sequence of amino acid residues from at or about 80 to at or about 155, inclusive of Figure 7 (SEQ ID NO:13), or (b) the complement of the DNA of (a).

In a still further aspect, the invention concerns an isolated nucleic acid molecule comprising (a) DNA encoding an IL-11p polypeptide having at least at or about 80% positives, or at least at or about 85% positives, or at least at or about 90% positives, or at least at or about 95% positives to an amino acid sequence selected from the group consisting of: (1) amino acid residues from at or about 2 to at or about 155, inclusive of Figure 7 (SEQ ID NO:13), and (2) amino acid residues from at or about 2 to at or about 155, inclusive of Figure 9 (SEQ ID NO:16), or (b) the complement of the DNA of (a).

In a still further aspect, the invention concerns an isolated nucleic acid molecule comprising (a) DNA encoding an IL-11p polypeptide that retains at least one biologic activity of a native sequence IL-11p, such as the IL-18R binding activity of a native sequence hIL-1Ra1, hIL-1Ra1L, or hIL-1Ra1V, and which IL-11p polypeptide has at least at or about 80% positives, or at least at or about 85% positives, or at least at or about 95% positives to an amino acid sequence selected from the group consisting of: (1) amino acid residues from at or about 1 to at or about 207, inclusive of Figure 15 (SEQ ID NO:19), and (2)

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amino acid residues from at or about 1 to at or about 218, inclusive of Figure 19 (SEQ ID NO:25), or (b) the complement of the DNA of (a).

In another embodiment, the invention provides a vector comprising DNA encoding IL-11p or its variants. The vector may comprise any of the isolated nucleic acid molecules hereinabove defined.

A host cell comprising such a vector is also provided. By way of example, the host cells may be CHO cells, *E. coli*, or yeast. A process for producing lL-1lp polypeptides is further provided and comprises culturing host cells under conditions suitable for expression of IL-1lp and recovering IL-1lp from the cell culture.

In another embodiment, the invention provides isolated IL-11p polypeptide encoded by any of the isolated nucleic acid sequences hereinabove defined.

In another aspect, the invention provides isolated native sequence IL-1lp polypeptide, which in one embodiment, comprises an amino acid sequence selected from the group consisting of: (1) the amino acid sequence of residues from at or about 37 to at or about 203, inclusive of Figure 2 (SEQ ID NO:5), (2) the amino acid sequence of residues from at or about 15 to at or about 193, inclusive of Figure 3 (SEQ ID NO:7), (3) the amino acid sequence of residues from at or about 34 to at or about 155, inclusive of Figure 7 (SEQ ID NO:13), and (4) the amino acid sequence of residues from at or about 34 to at or about 155, inclusive of Figure 9 (SEQ ID NO:16).

In another aspect, the invention provides isolated native sequence IL-1lp polypeptide, which in one embodiment, comprises an amino acid sequence selected from the group consisting of: (1) the amino acid sequence of residues from at or about 1 to at or about 207, inclusive of Figure 15 (SEQ ID NO:19), (2) the amino acid sequence of residues from at or about 26 to at or about 207, inclusive of Figure 15 (SEQ ID NO:19), (3) the amino acid sequence of residues from at or about 1 to at or about 167, inclusive of Figure 16 (SEQ ID NO:21), (4) the amino acid sequence of residues from at or about 167, inclusive of Figure 16 (SEQ ID NO:21), (5) the amino acid sequence of residues from at or about 1 to at or about 218, inclusive of Figure 19 (SEQ ID NO:25), (6) the amino acid sequence of residues from at or about 12 to at or about 218, inclusive of Figure 19 (SEQ ID NO:25), (7) the amino acid sequence of residues from at or about 37 to at or about 218, inclusive of Figure 19 (SEQ ID NO:25), and (8) the amino acid sequence of residues from at or about 218, inclusive of Figure 19 (SEQ ID NO:25).

In another aspect, the invention concerns an isolated IL-1lp polypeptide that retains at least one biologic activity of a native sequence IL-1lp, such as the IL-1R binding activity of a native sequence hIL-1Ra3 or mIL-1Ra3, or the IL-18R binding activity of a native sequence hIL-1Ra1L, or hIL-1Ra1V, and that consists of an amino acid sequence having at least at or about 80% sequence identity, or at least at or about 85% sequence identity, or at least at or about 90% sequence identity, or at least at or about 95% sequence identity to the sequence of amino acid residues from at or about 37 to at or about 203, inclusive of Figure 2 (SEQ ID NO:5), the sequence of amino acid residues from at or about 15 to at or about 193, inclusive of Figure 3 (SEQ ID NO:7), the sequence of amino acid residues from at or about 34 to at or about 155,

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inclusive of Figure 7 (SEQ ID NO:13), the sequence of amino acid residues from at or about 34 to at or about 155, inclusive of Figure 9 (SEQ ID NO:16), the sequence of amino acid residues from at or about 26 to at or about 207, inclusive of Figure 15 (SEQ ID NO:19), or the sequence of amino acid residues from at or about 46 to at or about 218, inclusive of Figure 19 (SEQ ID NO:25).

In another aspect, the invention concerns an isolated IL-1lp polypeptide that retains at least one biologic activity of a native sequence IL-1lp, such as the IL-18R binding activity of a native sequence hIL-1Ra1, hIL-1Ra1L, or hIL-1Ra1V, and that consists of an amino acid sequence having at least at or about 80% sequence identity, or at least at or about 85% sequence identity, or at least at or about 90% sequence identity, or at least at or about 95% sequence identity to the sequence of amino acid residues from at or about 1 to at or about 207, inclusive of Figure 15 (SEQ ID NO:19), or the sequence of amino acid residues from at or about 1 to at or about 218, inclusive of Figure 19 (SEO ID NO:25).

In another aspect, the invention provides an isolated IL-1lp selected from the group consisting of: (1) a polypeptide, such as an hIL-1Ra1 polypeptide, consisting of an amino acid sequence having a sequence identity of at least at or about 80%, or at least at or about 85%, or at least at or about 90%, or at least at or about 95%, to the sequence of amino acid residues from at or about 37 to at or about 63, inclusive of Figure 2 (SEQ ID NO:5); (2) a polypeptide, such as an hIL-1Ra1 polypeptide, consisting of an amino acid sequence having a sequence identity of at least at or about 80%, or at least at or about 85%, or at least at or about 90%, or at least at or about 95%, to the sequence of amino acid residues from at or about 15 to at or about 53, inclusive of Figure 3 (SEQ ID NO:7); (3) a polypeptide, such as an hIL-1Ra2 polypeptide, comprising the amino acid sequence of amino acid residues from at or about 1 to at or about 134, inclusive of Figure 5 (SEQ ID NO:10); (4) a polypeptide comprising the amino acid sequence of amino acid residues from at or about 10 to at or about 134, inclusive of Figure 5 (SEQ ID NO:10); (5) a polypeptide, such as an hIL-1Ra2 polypeptide, consisting of the amino acid sequence of amino acid residues from at or about 27 to at or about 134, inclusive of Figure 5 (SEQ ID NO:10); (6) a polypeptide, such as an hIL-1Ra2 fusion variant polypeptide, consisting of a native amino acid sequence of hIL-1Ra2 consisting of amino acid residues from at or about 27 to at or about 134, inclusive of Figure 5 (SEO ID NO:10) fused at its N-terminus or Cterminus to a heterologous amino acid or amino acid sequence; (7) a polypeptide, such as an hIL-1Ra3 polypeptide, consisting of an amino acid sequence having a sequence identity of at least at or about 80%, or at least at or about 85%, or at least at or about 90%, or at least at or about 95%, to the sequence of amino acid residues from at or about 95 to at or about 134, inclusive of Figure 7 (SEQ ID NO:13); and (8) a polypeptide, such as a mIL-1Ra3 polypeptide, consisting of an amino acid sequence having a sequence identity of at least at or about 80%, or at least at or about 85%, or at least at or about 90%, or at least at or about 95%, to the sequence of amino acid residues from at or about 95 to at or about 134, inclusive of Figure 9 (SEO ID NO:16).

In another aspect, the invention provides an isolated IL-1lp selected from the group consisting of: (1) a polypeptide, such as an hIL-1Ra1L polypeptide, consisting of an amino acid sequence having a sequence identity of at least at or about 80%, or at least at or about 85%, or

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at least at or about 90%, or at least at or about 95%, to the sequence of amino acid residues from at or about 27 to at or about 44, inclusive of Figure 15 (SEQ ID NO:19); (2) a polypeptide, such as an hIL-1Ra1S fusion variant polypeptide, consisting of a native amino acid sequence of hIL-1Ra1S consisting of amino acid residues from at or about 39 to at or about 167, inclusive of Figure 16 (SEQ ID NO:21) fused at its N-terminus or C-terminus to a heterologous amino acid or amino acid sequence; (3) a polypeptide, such as an hIL-1Ra1S fusion variant polypeptide, consisting of a native amino acid sequence of hIL-1Ra1S consisting of amino acid residues from at or about 47 to at or about 167, inclusive of Figure 16 (SEQ ID NO:21) fused at its N-terminus or C-terminus to a heterologous amino acid or amino acid sequence; (4) a polypeptide, such as an hIL-1Ra1S polypeptide, consisting of amino acid residues from at or about 39 to at or about 167, inclusive of Figure 16 (SEQ ID NO:21); (5) a polypeptide, such as an hIL-1Ra1S polypeptide, consisting of amino acid residues from at or about 47 to at or about 167, inclusive of Figure 16 (SEQ ID NO:21); and (6) a polypeptide, such as an hIL-1Ra1V polypeptide, consisting of an amino acid sequence having a sequence identity of at least at or about 80%, or at least at or about 85%, or at least at or about 90%, or at least at or about 95%, to the sequence of amino acid residues from at or about 37 to at or about 55, inclusive of Figure 19 (SEQ ID NO:25).

In another aspect, the invention concerns an isolated IL-11p polypeptide that consists of an amino acid sequence having at least at or about 80% sequence identity, or at least at or about 85% sequence identity, or at least at or about 95% sequence identity to the sequence of amino acid residues from at or about 37 to at or about 203, inclusive of Figure 2 (SEQ ID NO:5), the sequence of amino acid residues from at or about 195 to at or about 193, inclusive of Figure 3 (SEQ ID NO:7), the sequence of amino acid residues from at or about 155, inclusive of Figure 7 (SEQ ID NO:13), the sequence of amino acid residues from at or about 34 to at or about 34 to at or about 34 to at or about 255, inclusive of Figure 9 (SEQ ID NO:16), the sequence of amino acid residues from at or about 26 to at or about 207, inclusive of Figure 15 (SEQ ID NO:19), or the sequence of amino acid residues from at or about 218, inclusive of Figure 19 (SEQ ID NO:25).

In another aspect, the invention concerns an isolated IL-1lp polypeptide that consists of an amino acid sequence having at least at or about 80% sequence identity, or at least at or about 85% sequence identity, or at least at or about 90% sequence identity, or at least at or about 95% sequence identity to the sequence of amino acid residues from at or about 1 to at or about 207, inclusive of Figure 15 (SEQ ID NO:19), or the sequence of amino acid residues from at or about 1 to at or about 218, inclusive of Figure 19 (SEQ ID NO:25).

In another aspect, the invention provides an isolated polypeptide, such as an hIL-1Ra3 polypeptide, consisting of an amino acid sequence having a sequence identity of at least at or about 80%, or at least at or about 85%, or at least at or about 90%, or at least at or about 95%, to the sequence of amino acid residues from at or about 80 to at or about 155, inclusive of Figure 7 (SEQ ID NO:13).

In a further aspect, the invention concerns an isolated IL-11p polypeptide that retains at least one biologic activity of a native sequence IL-11p, such as the IL-1R binding activity of a native sequence hIL-1Ra3 or mIL-1Ra3, or the IL-18R binding activity of a native sequence hIL-

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1Ra1, hIL-1Ra1L, or hIL-1Ra1V, and that consists of an amino acid sequence having at least at or about 80% sequence identity, or at least at or about 85% sequence identity, or at least at or about 90% sequence identity, or at least at or about 95% sequence identity to the amino acid sequence of an IL-1lp, such as a mature IL-1lp polypeptide, encoded by the cDNA insert in the vector deposited as ATCC Deposit No. 203588 (DNA85066-2534), ATCC Deposit No. 203587 (DNA96786-2534), ATCC Deposit No. 203589 (DNA96787-2534), ATCC Deposit No. 203590 (DNA92505-2534), ATCC Deposit No. 203590 (DNA92505-2534), ATCC Deposit No. 203590 (DNA114876-2534). In a preferred embodiment, the IL-1lp polypeptide comprises the amino acid sequence of an IL-1lp, such as a mature IL-1lp polypeptide, encoded by the cDNA insert in the vector deposited as ATCC Deposit No. 203588 (DNA85066-2534), ATCC Deposit No. 203587 (DNA96786-2534), ATCC Deposit No. 203586 (DNA92929-2534), ATCC Deposit No. 203589 (DNA96787-2534), ATCC Deposit No. 203590 (DNA92505-2534), ATCC Deposit No. 203846 (DNA102043-2534), ATCC Deposit No. 203973 (DNA114876-2534), or ATCC Deposit No. 203855 (DNA102043-2534), ATCC Deposit No. 203973 (DNA114876-2534), or ATCC Deposit No. 203855 (DNA102043-2534), ATCC Deposit No. 203973 (DNA114876-2534), or ATCC Deposit No. 203855 (DNA102043-2534), ATCC Deposit No. 203973 (DNA114876-2534), or ATCC Deposit No. 203855 (DNA102043-2534), ATCC Deposit No. 203973 (DNA114876-2534), or ATCC Deposit No. 203855 (DNA102043-2534)

In a further aspect, the invention concerns an isolated IL-1lp polypeptide that retains at least one biologic activity of a native sequence IL-11p, such as the IL-1R binding activity of a native sequence hIL-1Ra3 or mIL-1Ra3, or the IL-18R binding activity of a native sequence hIL-1Ra1, hIL-1Ra1L, or hIL-1Ra1V, and that consists of an amino acid sequence having at least at or about 80% sequence identity, or at least at or about 85% sequence identity, or at least at or about 90% sequence identity, or at least at or about 95% sequence identity to the entire amino acid sequence encoded by the longest open reading frame in the cDNA insert of a vector selected from the group consisting of the vectors deposited as ATCC Deposit No. 203588 (DNA85066-2534), ATCC Deposit No. 203587 (DNA96786-2534), ATCC Deposit No. 203589 (DNA96787-2534), ATCC Deposit No. 203590 (DNA92505-2534), ATCC Deposit No. 203846 (DNA102043-2534), and ATCC Deposit No. 203973 (DNA114876-2534). In a preferred embodiment, the IL-1lp polypeptide comprises the entire amino acid sequence encoded by the longest open reading frame in the cDNA insert of a vector selected from the group consisting of the vectors deposited as ATCC Deposit No. 203588 (DNA85066-2534), ATCC Deposit No. 203587 (DNA96786-2534), ATCC Deposit No. 203586 (DNA92929-2534), ATCC Deposit No. 203589 (DNA96787-2534), and ATCC Deposit No. 203590 (DNA92505-2534). In another preferred embodiment, the IL-1lp polypeptide comprises the entire amino acid sequence encoded by the longest open reading frame in the cDNA insert of a vector selected from the group consisting of the vectors deposited as ATCC Deposit No. 203846 (DNA102043-2534), ATCC Deposit No. 203855 (DNA102044-2534), and ATCC Deposit No. 203973 (DNA114876-2534).

In another aspect, the invention concerns an isolated IL-11p polypeptide that retains at least one biologic activity of a native sequence IL-11p, such as the IL-1R binding activity of a native sequence IL-1Ra3 or mIL-1Ra3, or the IL-18b binding activity of a native sequence hIL-1Ra1, hIL-1Ra1L, or hIL-1Ra1V, and that consists of an amino acid sequence having at least at or about 80% sequence identity, or at least at or about 85% sequence identity, or at least at or about 95% sequence identity to an amino acid sequence selected from the group consisting of: (1) the entire amino acid sequence encoded by

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the longest open reading frame in the cDNA insert in the vector deposited as ATCC Deposit No. 203588, (2) the entire amino acid sequence, or the entire amino acid sequence excluding the 36 N-terminal amino acid residues of such sequence, encoded by the longest open reading frame in the cDNA insert in the vector deposited as ATCC Deposit No. 203587, (3) the entire amino acid sequence, or the entire amino acid sequence excluding the N-terminal amino acid residue of such sequence, encoded by the longest open reading frame in the cDNA insert in the vector deposited as ATCC Deposit No. 203589, (4) the entire amino acid sequence, or the entire amino acid sequence excluding the N-terminal amino acid residue of such sequence, encoded by the longest open reading frame in the cDNA insert in the vector deposited as ATCC Deposit No. 203590, (5) the entire amino acid sequence, or the entire amino acid sequence excluding the N-terminal amino acid residue of such sequence, or the entire amino acid sequence excluding the 34 Nterminal amino acid residues of such sequence, encoded by the longest open reading frame in the cDNA insert in the vector deposited as ATCC Deposit No. 203846, and (6) the entire amino acid sequence, or the entire amino acid sequence excluding the N-terminal amino acid residue of such sequence, or the entire amino acid sequence excluding the 45 N-terminal amino acid residues of such sequence, encoded by the longest open reading frame in the cDNA insert in the vector deposited as ATCC Deposit No. 203973.

In a further aspect, the invention concerns an isolated IL-1lp polypeptide that consists of an amino acid sequence having at least at or about 80% sequence identity, or at least at or about 85% sequence identity, or at least at or about 95% sequence identity to the amino acid sequence of an IL-1lp, such as a mature IL-1lp polypeptide, encoded by the cDNA insert in the vector deposited as ATCC Deposit No. 203588 (DNA85066-2534), ATCC Deposit No. 203589 (DNA96787-2534), ATCC Deposit No. 203589 (DNA92505-2534), ATCC Deposit No. 203846 (DNA102043-2534), or ATCC Deposit No. 203973 (DNA114876-2534).

In a further aspect, the invention concerns an isolated IL-1lp polypeptide that consists of an amino acid sequence having at least at or about 80% sequence identity, or at least at or about 85% sequence identity, or at least at or about 95% sequence identity, or at least at or about 95% sequence identity to the entire amino acid sequence encoded by the longest open reading frame in the cDNA insert of a vector selected from the group consisting of the vectors deposited as ATCC Deposit No. 203588 (DNA85066-2534), ATCC Deposit No. 203587 (DNA96786-2534), ATCC Deposit No. 203590 (DNA96786-2534), ATCC Deposit No. 203846 (DNA102043-2534), and ATCC Deposit No. 203973 (DNA114876-2534).

In another aspect, the invention concerns an isolated IL-1lp polypeptide that consists of an amino acid sequence having at least at or about 80% sequence identity, or at least at or about 95% sequence identity, or at least at or about 95% sequence identity to an amino acid sequence selected from the group consisting of: (1) the entire amino acid sequence encoded by the longest open reading frame in the cDNA insert in the vector deposited as ATCC Deposit No. 203588, (2) the entire amino acid sequence, or the entire amino acid sequence excluding the 36 N-terminal amino acid residues of such sequence, encoded by

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the longest open reading frame in the cDNA insert in the vector deposited as ATCC Deposit No. 203587, (3) the entire amino acid sequence, or the entire amino acid sequence excluding the N-terminal amino acid residue of such sequence, encoded by the longest open reading frame in the cDNA insert in the vector deposited as ATCC Deposit No. 203589, (4) the entire amino acid sequence, or the entire amino acid sequence excluding the N-terminal amino acid residue of such sequence, encoded by the longest open reading frame in the cDNA insert in the vector deposited as ATCC Deposit No. 203590, (5) the entire amino acid sequence, or the entire amino acid sequence excluding the N-terminal amino acid residue of such sequence, or the entire amino acid sequence excluding the 34 N-terminal amino acid residues of such sequence, encoded by the longest open reading frame in the cDNA insert in the vector deposited as ATCC Deposit No. 203846, and (6) the entire amino acid sequence, or the entire amino acid sequence excluding the N-terminal amino acid residue of such sequence, or the entire amino acid sequence excluding the 45 N-terminal amino acid residues of such sequence, encoded by the longest open reading frame in the cDNA insert in the vector deposited as ATCC Deposit No. 203973.

In a preferred embodiment, the IL-11p polypeptide comprises an amino acid sequence selected from the group consisting of: (1) the entire amino acid sequence encoded by the longest open reading frame in the cDNA insert in the vector deposited as ATCC Deposit No. 203588, (2) the entire amino acid sequence, or the entire amino acid sequence excluding the 36 N-terminal amino acid residues of such sequence, encoded by the longest open reading frame in the cDNA insert in the vector deposited as ATCC Deposit No. 203587, (3) the entire amino acid sequence, or the entire amino acid sequence excluding the N-terminal amino acid residue of such sequence, encoded by the longest open reading frame in the cDNA insert in the vector deposited as ATCC Deposit No. 203589, (4) the entire amino acid sequence, or the entire amino acid sequence excluding the N-terminal amino acid residue of such sequence, encoded by the longest open reading frame in the cDNA insert in the vector deposited as ATCC Deposit No. 203590, (5) the entire amino acid sequence, or the entire amino acid sequence excluding the N-terminal amino acid residue of such sequence, or the entire amino acid sequence excluding the 34 N-terminal amino acid residues of such sequence, encoded by the longest open reading frame in the cDNA insert in the vector deposited as ATCC Deposit No. 203846, and (6) the entire amino acid sequence, or the entire amino acid sequence excluding the N-terminal amino acid residue of such sequence, or the entire amino acid sequence excluding the 45 N-terminal amino acid residues of such sequence, encoded by the longest open reading frame in the cDNA insert in the vector deposited as ATCC Deposit No. 203973.

In another aspect, the invention concerns an isolated IL-1Ip polypeptide that retains at least one biologic activity of a native sequence IL-1Ip, such as the IL-1R binding activity of a native sequence hIL-1Ra3 or mIL-1Ra3, or the IL-18R binding activity of a native sequence hIL-1Ra1, hIL-1Ra1L, or hIL-1Ra1V, and that consists of an amino acid sequence having at least at or about 80% positives, or at least at or about 85% positives, or at least at or about 95% positives to the sequence of amino acid residues from at or about 37 to at or about 203, inclusive of Figure 2 (SEQ ID NO:5), the sequence of amino acid residues from at or about 15 to at or about 193, inclusive of Figure 3 (SEQ ID NO:7), the

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sequence of amino acid residues from at or about 34 to at or about 155, inclusive of Figure 7 (SEQ ID NO:13), the sequence of amino acid residues from at or about 34 to at or about 155, inclusive of Figure 9 (SEQ ID NO:16), or the sequence of amino acid residues from at or about 46 to at or about 218, inclusive of Figure 19 (SEQ ID NO:25).

In another aspect, the invention concerns an isolated IL-1lp polypeptide that retains at least one biologic activity of a native sequence IL-1lp, such as the IL-18R binding activity of a native sequence hIL-1Ra1, hIL-1Ra1L, or hIL-1Ra1V, and that consists of an amino acid sequence having at least at or about 80% positives, or at least at or about 85% positives, or at least at or about 90% positives, or at least at or about 95% positives to the sequence of amino acid residues from at or about 1 to at or about 207, inclusive of Figure 15 (SEQ ID NO:19), or the sequence of amino acid residues from at or about 1 to at or about 218, inclusive of Figure 19 (SEO ID NO:25).

In another aspect, the invention provides an isolated IL-1lp selected from the group consisting of: (1) a polypeptide, such as an hIL-1Ra1 polypeptide, consisting of an amino acid sequence having a % positives value of at least at or about 80%, or at least at or about 85%, or at least at or about 90%, or at least at or about 95%, to the sequence of amino acid residues from at or about 37 to at or about 63, inclusive of Figure 2 (SEQ ID NO:5); (2) a polypeptide, such as an hIL-1Ra1 polypeptide, consisting of an amino acid sequence having a % positives value of at least at or about 80%, or at least at or about 85%, or at least at or about 90%, or at least at or about 95%, to the sequence of amino acid residues from at or about 15 to at or about 53, inclusive of Figure 3 (SEQ ID NO:7); (3) a polypeptide, such as an hlL-1Ra2 polypeptide, comprising the amino acid sequence of amino acid residues from at or about 1 to at or about 134, inclusive of Figure 5 (SEQ ID NO:10); (4) a polypeptide comprising the amino acid sequence of amino acid residues from at or about 10 to at or about 134, inclusive of Figure 5 (SEQ ID NO:10); (5) a polypeptide, such as an hIL-1Ra2 polypeptide, consisting of the amino acid sequence of amino acid residues from at or about 27 to at or about 134, inclusive of Figure 5 (SEO ID NO:10); (6) a polypeptide, such as an hIL-1Ra2 fusion variant polypeptide, consisting of a native amino acid sequence of hIL-1Ra2 consisting of amino acid residues from at or about 27 to at or about 134, inclusive of Figure 5 (SEQ ID NO:10) fused at its N-terminus or Cterminus to a heterologous amino acid or amino acid sequence; (7) a polypeptide, such as an hIL-1Ra3 polypeptide, consisting of an amino acid sequence having a % positives value of at least at or about 80%, or at least at or about 85%, or at least at or about 90%, or at least at or about 95%, to the sequence of amino acid residues from at or about 95 to at or about 134, inclusive of Figure 7 (SEQ ID NO:13); and (8) a polypeptide, such as a mIL-1Ra3 polypeptide, consisting of an amino acid sequence having a % positives value of at least at or about 80%, or at least at or about 85%, or at least at or about 90%, or at least at or about 95%, to the sequence of amino acid residues from at or about 95 to at or about 134, inclusive of Figure 9 (SEQ ID NO:16).

In another aspect, the invention provides an isolated IL-1lp selected from the group consisting of: (1) a polypeptide, such as an hIL-1Ra1L polypeptide, consisting of an amino acid sequence having a % positives value of at least at or about 80%, or at least at or about 95%, to the sequence of amino acid residues

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from at or about 27 to at or about 44, inclusive of Figure 15 (SEQ ID NO:19); (2) a polypeptide, such as an hIL-1Ra1S fusion variant polypeptide, consisting of a native amino acid sequence of hIL-1Ra1S consisting of amino acid residues from at or about 39 to at or about 167, inclusive of Figure 16 (SEQ ID NO:21) fused at its N-terminus or C-terminus to a heterologous amino acid or amino acid sequence; (3) a polypeptide, such as an hIL-1Ra1S fusion variant polypeptide, consisting of a native amino acid sequence of hIL-1Ra1S consisting of amino acid residues from at or about 47 to at or about 167, inclusive of Figure 16 (SEQ ID NO:21) fused at its N-terminus or C-terminus to a heterologous amino acid or amino acid sequence; (4) a polypeptide, such as an hIL-1Ra1S polypeptide, consisting of amino acid residues from at or about 39 to at or about 167, inclusive of Figure 16 (SEQ ID NO:21); (5) a polypeptide, such as an hIL-1Ra1S polypeptide, consisting of amino acid residues from at or about 167, inclusive of Figure 16 (SEQ ID NO:21); and (6) a polypeptide, such as an hIL-1Ra1V polypeptide, consisting of amino acid sequence having a % positives value of at least at or about 80%, or at least at or about 90%, or at least at or about 90%, or at least at or about 37 to at or about 55, inclusive of Figure 19 (SEQ ID NO:25).

In another aspect, the invention concerns an isolated IL-1lp polypeptide that consists of an amino acid sequence having at least at or about 80% positives, or at least at or about 85% positives, or at least at or about 90% positives, or at least at or about 95% positives to the sequence of amino acid residues from at or about 37 to at or about 203, inclusive of Figure 2 (SEQ ID NO:5), the sequence of amino acid residues from at or about 15 to at or about 193, inclusive of Figure 3 (SEQ ID NO:7), the sequence of amino acid residues from at or about 194, inclusive of Figure 5 (SEQ ID NO:13), the sequence of amino acid residues from at or about 34 to at or about 155, inclusive of Figure 9 (SEQ ID NO:16), or the sequence of amino acid residues from at or about 46 to at or about 218, inclusive of Figure 19 (SEQ ID NO:25).

In another aspect, the invention concerns an isolated IL-1lp polypeptide that consists of an amino acid sequence having at least at or about 80% positives, or at least at or about 85% positives, or at least at or about 95% positives to the sequence of amino acid residues from at or about 1 to at or about 207, inclusive of Figure 15 (SEQ ID NO:19), or the sequence of amino acid residues from at or about 1 to at or about 218, inclusive of Figure 19 (SEQ ID NO:25).

In a further aspect, the invention concerns an isolated IL-1lp polypeptide that retains at least one biologic activity of a native sequence IL-1lp, such as the IL-1R binding activity of a native sequence hIL-1Ra3, or mIL-1Ra3, or the IL-18R binding activity of a native sequence hIL-1Ra1, hIL-1Ra1L, or hIL-1Ra1V, and that consists of an amino acid sequence that has at least at or about 80% positives, or at least at or about 85% positives, or at least at or about 95% positives to the amino acid sequence of an IL-1lp, such as a mature IL-1lp polypeptide, encoded by the cDNA insert in the vector deposited as ATCC Deposit No. 203588 (DNA95066-2534), ATCC Deposit No. 203589 (DNA96787-2534), ATCC Deposit No. 203599 (DNA95786-2534), ATCC Deposit No. 203593 (DNA95786-2534), ATCC Deposit No. 203846 (DNA102043-2534), or ATCC Deposit No. 203973 (DNA114876-2534).

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In a further aspect, the invention concerns an isolated IL-Ilp polypeptide that retains at least one biologic activity of a native sequence IL-Ilp, such as the IL-IR binding activity of a native sequence hIL-IRa3 or mIL-IRa3, or the IL-IRR binding activity of a native sequence hIL-IRa1, hIL-IRa1L, or hIL-IRa1V, and that consists of an amino acid sequence that has at least at or about 80% positives, or at least at or about 95% positives, or at least at or about 95% positives, or at least at or about 95% positives to the entire amino acid sequence encoded by the longest open reading frame in the cDNA insert of a vector selected from the group consisting of the vectors deposited as ATCC Deposit No. 203588 (DNA85066-2534), ATCC Deposit No. 203587 (DNA96786-2534), ATCC Deposit No. 203590 (DNA92505-2534), ATCC Deposit No. 203590 (DNA92505-2534), ATCC Deposit No. 203973 (DNA14876-2534), ATCC Deposit No. 203973 (DNA14876-2534).

In another aspect, the invention concerns an isolated IL-1lp polypeptide that retains at least one biologic activity of a native sequence IL-1lp, such as the IL-1R binding activity of a native sequence IL-1Ra3 or mIL-1Ra3, or the IL-18R binding activity of a native sequence hIL-1Ra1, hIL-1Ra1L, or hIL-1Ra1V, and that consists of an amino acid sequence having at least at or about 80% positives, or at least at or about 85% positives, or at least at or about 90% positives, or at least at or about 95% positives to an amino acid sequence selected from the group consisting of: (1) the entire amino acid sequence encoded by the longest open reading frame in the cDNA insert in the vector deposited as ATCC Deposit No. 203588, (2) the entire amino acid sequence, or the entire amino acid sequence excluding the 36 N-terminal amino acid residues of such sequence, encoded by the longest open reading frame in the cDNA insert in the vector deposited as ATCC Deposit No. 203587, (3) the entire amino acid sequence, or the entire amino acid sequence excluding the N-terminal amino acid residue of such sequence, encoded by the longest open reading frame in the cDNA insert in the vector deposited as ATCC Deposit No. 203589, (4) the entire amino acid sequence, or the entire amino acid sequence excluding the N-terminal amino acid residue of such sequence, encoded by the longest open reading frame in the cDNA insert in the vector deposited as ATCC Deposit No. 203590, (5) the entire amino acid sequence, or the entire amino acid sequence excluding the N-terminal amino acid residue of such sequence, or the entire amino acid sequence excluding the 34 N-terminal amino acid residues of such sequence, encoded by the longest open reading frame in the cDNA insert in the vector deposited as ATCC Deposit No. 203846, and (6) the entire amino acid sequence, or the entire amino acid sequence excluding the N-terminal amino acid residue of such sequence, or the entire amino acid sequence excluding the 45 N-terminal amino acid residues of such sequence, encoded by the longest open reading frame in the cDNA insert in the vector deposited as ATCC Deposit No. 203973.

In a further aspect, the invention concerns an isolated IL-1lp polypeptide that consists of an amino acid sequence that has at least at or about 80% positives, or at least at or about 85% positives, or at least at or about 90% positives, or at least at or about 95% positives to the amino acid sequence of an IL-1lp, such as a mature IL-1lp polypeptide, encoded by the cDNA insert in the vector deposited as ATCC Deposit No. 203588 (DNA85066-2534), ATCC Deposit No. 203587 (DNA96786-2534), ATCC Deposit No.

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203590 (DNA92505-2534), ATCC Deposit No. 203846 (DNA102043-2534), or ATCC Deposit No. 203973 (DNA114876-2534).

In a further aspect, the invention concerns an isolated IL-1lp polypeptide that consists of an amino acid sequence that has at least at or about 80% positives, or at least at or about 85% positives, or at least at or about 90% positives, or at least at or about 95% positives to the entire amino acid sequence encoded by the longest open reading frame in the cDNA insert of a vector selected from the group consisting of the vectors deposited as ATCC Deposit No. 203588 (DNA85066-2534), ATCC Deposit No. 203587 (DNA96786-2534), ATCC Deposit No. 203589 (DNA96787-2534), ATCC Deposit No. 203580 (DNA92505-2534), ATCC Deposit No. 203846 (DNA102043-2534), and ATCC Deposit No. 203973 (DNA114876-2534).

In another aspect, the invention concerns an isolated IL-1lp polypeptide that consists of an amino acid sequence having at least at or about 80% positives, or at least at or about 85% positives, or at least at or about 90% positives, or at least at or about 95% positives to an amino acid sequence selected from the group consisting of: (1) the entire amino acid sequence encoded by the longest open reading frame in the cDNA insert in the vector deposited as ATCC Deposit No. 203588, (2) the entire amino acid sequence, or the entire amino acid sequence excluding the 36 N-terminal amino acid residues of such sequence, encoded by the longest open reading frame in the cDNA insert in the vector deposited as ATCC Deposit No. 203587, (3) the entire amino acid sequence, or the entire amino acid sequence excluding the N-terminal amino acid residue of such sequence, encoded by the longest open reading frame in the cDNA insert in the vector deposited as ATCC Deposit No. 203589, (4) the entire amino acid sequence, or the entire amino acid sequence excluding the N-terminal amino acid residue of such sequence, encoded by the longest open reading frame in the cDNA insert in the vector deposited as ATCC Deposit No. 203590, (5) the entire amino acid sequence, or the entire amino acid sequence excluding the N-terminal amino acid residue of such sequence, or the entire amino acid sequence excluding the 34 Nterminal amino acid residues of such sequence, encoded by the longest open reading frame in the cDNA insert in the vector deposited as ATCC Deposit No. 203846, and (6) the entire amino acid sequence, or the entire amino acid sequence excluding the N-terminal amino acid residue of such sequence, or the entire amino acid sequence excluding the 45 N-terminal amino acid residues of such sequence, encoded by the longest open reading frame in the cDNA insert in the vector deposited as ATCC Deposit No. 203973.

In another aspect, the invention provides an isolated polypeptide, such as an hIL-1Ra3 polypeptide, consisting of an amino acid sequence having a % positives value of at least at or about 80%, or at least at or about 85%, or at least at or about 90%, or at least at or about 95%, to the sequence of amino acid residues from at or about 80 to at or about 155, inclusive of Figure 7 (SEQ ID NO:13).

In yet another aspect, the invention concerns an isolated IL-11p polypeptide comprising an amino acid sequence selected from the group consisting of: (1) amino acid residues from at or about 37 to at or about 63, inclusive of Figure 2 (SEQ ID NO:5); (2) amino acid residues from at or about 15 to at or about 53, inclusive of Figure 3 (SEQ ID NO:7); (3) amino acid residues from at or about 80 to at or about 155, inclusive of Figure 7 (SEQ ID NO:13); and (4) amino acid

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residues from at or about 95 to at or about 155, inclusive of Figure 9 (SEQ ID NO:16), or a fragment of such IL-11p polypeptide that coincides with a stretch of at least about 10 contiguous amino acids in such amino acid sequence, wherein the IL-11p polypeptide or fragment thereof is sufficient to provide a binding site for an anti-IL-11p antibody. Preferably, the IL-11p fragment retains at least one biologic activity of a native sequence IL-11p polypeptide.

In yet another aspect, the invention concerns an isolated IL-1lp polypeptide comprising an amino acid sequence selected from the group consisting of: (1) amino acid residues from at or about 26 to at or about 44, inclusive of Figure 15 (SEQ ID NO:19), (2) amino acid residues from at or about 26 to at or about 78, inclusive of Figure 19 (SEQ ID NO:25), and (3) amino acid residues from at or about 46 to at or about 89, inclusive of Figure 19 (SEQ ID NO:25), or a fragment of such IL-1lp polypeptide that coincides with a stretch of at least about 10 contiguous amino acids in such amino acid sequence, wherein the IL-1lp polypeptide or fragment thereof is sufficient to provide a binding site for an anti-IL-1lp antibody. Preferably, the IL-1lp fragment retains at least one biologic activity of a native sequence IL-1lp polypeptide.

In a further aspect, the invention provides an isolated IL-1lp polypeptide selected from the group consisting of: (1) a polypeptide, such as an hIL-1Ra1 polypeptide, comprising amino acid residues from at or about 37 to at or about 63, inclusive of Figure 2 (SEQ ID NO:5); (2) a polypeptide, such as an hIL-1Ra1 polypeptide, comprising amino acid residues from at or about 15 to at or about 53, inclusive of Figure 3 (SEQ ID NO:7); (3) a polypeptide, such as an hIL-1Ra3 polypeptide, comprising amino acid residues from at or about 95 to at or about 134, inclusive of Figure 7 (SEQ ID NO:13); and (4) a polypeptide, such as a mIL-1Ra3 polypeptide, comprising amino acid residues from at or about 134, inclusive of Figure 9 (SEQ ID NO:16).

In a further aspect, the invention provides an isolated IL-1lp polypeptide selected from the group consisting of: (1) a polypeptide, such as an hIL-1Ra1 polypeptide, comprising amino acid residues from at or about 37 to at or about 203, inclusive of Figure 2 (SEQ ID NO:5); (2) a polypeptide, such as an hIL-1Ra1 polypeptide, comprising amino acid residues from at or about 15 to at or about 193, inclusive of Figure 3 (SEQ ID NO:7); (3) a polypeptide, such as an hIL-1Ra3 polypeptide, comprising amino acid residues from at or about 34 to at or about 155, inclusive of Figure 7 (SEQ ID NO:13); and (3) a polypeptide, such as a mIL-1Ra3 polypeptide, comprising amino acid residues from at or about 34 to at or about 155, inclusive of Figure 9 (SEQ ID NO:16).

In a further aspect, the invention provides an isolated polypeptide, such as an hIL-1Ra3 polypeptide, comprising amino acid residues from at or about 80 to at or about 155, inclusive of Figure 7 (SEQ ID NO:13).

In a further aspect, the invention provides an isolated IL-1lp polypeptide selected from the group consisting of: (1) a polypeptide, such as an hIL-1Ra1L polypeptide, comprising amino acid residues from at or about 26 to at or about 44, inclusive of Figure 15 (SEQ ID NO:19); (2) a polypeptide, such as an hIL-1Ra1L polypeptide, comprising amino acid residues from at or about 1 to at or about 44, inclusive of Figure 15 (SEQ ID NO:19); (3) a polypeptide, such as an hIL-1Ra1S polypeptide, comprising amino acid residues from at or about 1 to at or about 38, inclusive of Figure 16 (SEQ ID NO:21); (4) a polypeptide, such as an hIL-1Ra1V polypeptide,

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comprising amino acid residues from at or about 37 to at or about 55, inclusive of Figure 19 (SEQ ID NO:25); (5) a polypeptide, such as an hIL-1Ra1V polypeptide, comprising amino acid residues from at or about 12 to at or about 55, inclusive of Figure 19 (SEQ ID NO:25); and (6) a polypeptide, such as an hIL-1Ra1V polypeptide, comprising amino acid residues from at or about 1 to at or about 55, inclusive of Figure 19 (SEQ ID NO:25).

In a further aspect, the invention provides an isolated IL-11p polypeptide selected from the group consisting of: (1) a polypeptide, such as an hIL-1Ra1L fusion variant polypeptide, consisting of a native amino acid sequence of hIL-1Ra1L consisting of amino acid residues from at or about 1 to at or about 207, inclusive of Figure 15 (SEQ ID NO:19) fused at its N-terminus or C-terminus to a heterologous amino acid or amino acid sequence; (2) a polypeptide, such as an hIL-1Ra1L fusion variant polypeptide, consisting of a native amino acid sequence of hIL-1Ra1L consisting of amino acid residues from at or about 26 to at or about 207, inclusive of Figure 15 (SEQ ID NO:19) fused at its N-terminus or C-terminus to a heterologous amino acid or amino acid sequence; (3) a polypeptide, such as an hIL-1Ra1L polypeptide, consisting of amino acid residues from at or about 1 to at or about 207, inclusive of Figure 15 (SEQ ID NO:19); (4) a polypeptide, such as an hIL-1Ra1L polypeptide, consisting of amino acid residues from at or about 26 to at or about 207, inclusive of Figure 15 (SEQ ID NO:19); (5) a polypeptide, such as an hIL-1Ra1V fusion variant polypeptide, consisting of a native amino acid sequence of hIL-1Ra1V consisting of amino acid residues from at or about 1 to at or about 218, inclusive of Figure 19 (SEQ ID NO:25) fused at its N-terminus or C-terminus to a heterologous amino acid or amino acid sequence; (6) a polypeptide, such as an hIL-1Ra1V fusion variant polypeptide, consisting of a native amino acid sequence of hIL-1Ra1V consisting of amino acid residues from at or about 12 to at or about 218, inclusive of Figure 19 (SEQ ID NO:25) fused at its N-terminus or C-terminus to a heterologous amino acid or amino acid sequence; (7) a polypeptide, such as an hIL-1Ra1V fusion variant polypeptide, consisting of a native amino acid sequence of hIL-1Ra1V consisting of amino acid residues from at or about 37 to at or about 218, inclusive of Figure 19 (SEQ ID NO:25) fused at its N-terminus or C-terminus to a heterologous amino acid or amino acid sequence; (8) a polypeptide, such as an hIL-1Ra1V fusion variant polypeptide, consisting of a native amino acid sequence of hIL-1Ra1V consisting of amino acid residues from at or about 46 to at or about 218, inclusive of Figure 19 (SEQ ID NO:25) fused at its N-terminus or C-terminus to a heterologous amino acid or amino acid sequence; (9) a polypeptide, such as an hIL-1Ra1V polypeptide, consisting of the amino acid sequence of amino acid residues from at or about 1 to at or about 218, inclusive of Figure 19 (SEQ ID NO:25); (10) a polypeptide, such as an hIL-1Ra1V polypeptide, consisting of the amino acid sequence of amino acid residues from at or about 12 to at or about 218, inclusive of Figure 19 (SEQ ID NO:25); (11) a polypeptide, such as an hIL-1Ra1V polypeptide, consisting of the amino acid sequence of amino acid residues from at or about 37 to at or about 218, inclusive of Figure 19 (SEQ ID NO:25); and (12) a polypeptide, such as an hIL-1Ra1V polypeptide, consisting of the amino acid sequence of amino acid residues from at or about 46 to at or about 218, inclusive of Figure 19 (SEQ ID NO:25).

In a further aspect, the invention provides an isolated IL-11p polypeptide selected from the group consisting of: (1) a polypeptide, such as an hIL-1Ra1L polypeptide, comprising amino

PCT/US99/30720 WO 00/39297

acid residues from at or about 1 to at or about 207, inclusive of Figure 15 (SEQ ID NO:19); (2) a polypeptide, such as an hIL-1Ra1L polypeptide, comprising amino acid residues from at or about 26 to at or about 207, inclusive of Figure 15 (SEQ ID NO:19); (3) a polypeptide, such as an hIL-1Ra1S polypeptide, comprising amino acid residues from at or about 1 to at or about 167, inclusive of Figure 16 (SEQ ID NO:21); (4) a polypeptide, such as an hIL-1Ra1S polypeptide, comprising amino acid residues from at or about 26 to at or about 167, inclusive of Figure 16 (SEQ ID NO:21); (5) a polypeptide, such as an hIL-1Ra1V polypeptide, comprising amino acid residues from at or about 1 to at or about 218, inclusive of Figure 19 (SEQ ID NO:25); and (6) a polypeptide, such as an hIL-1RalV polypeptide, comprising amino acid residues from at or about 12 to at or about 218, inclusive of Figure 19 (SEO ID NO:25).

In a further aspect, the invention provides an isolated IL-1lp polypeptide selected from the group consisting of: (1) a polypeptide, such as an hIL-1Ra1V polypeptide, comprising amino acid residues from at or about 37 to at or about 218, inclusive of Figure 19 (SEQ ID NO:25); and (2) a polypeptide, such as an hIL-1Ra1V polypeptide, comprising amino acid residues from at or about 46 to at or about 218, inclusive of Figure 19 (SEQ ID NO:25).

In a further aspect, the invention provides an isolated IL-1lp polypeptide selected from the group consisting of: (1) a polypeptide consisting of the amino acid sequence of amino acid residues 10 to 134, inclusive of Figure 5 (SEO ID NO:10) fused at its N-terminus or C-terminus to a heterologous amino acid or amino acid sequence; (2) a polypeptide consisting of the amino acid sequence of amino acid residues 10 to 134, inclusive of Figure 5 (SEQ ID NO:10); (3) a polypeptide, such as an hIL-1Ra3 polypeptide, consisting of a native amino acid sequence of hIL-1Ra3 consisting of amino acid residues 2 to 155, inclusive of Figure 7 (SEQ ID NO:13) fused at its N-terminus or C-terminus to a heterologous amino acid or amino acid sequence; (4) a polypeptide, such as an hIL-1Ra3 polypeptide, consisting of the amino acid sequence of amino acid residues from 2 to 155, inclusive of Figure 7 (SEQ ID NO:13); (5) a polypeptide, such as a mIL-1Ra3 polypeptide, consisting of a native amino acid sequence of mIL-1Ra3 consisting of amino acid residues 2 to 155, inclusive of Figure 9 (SEQ ID NO:16) fused at its N-terminus or C-terminus to a heterologous amino acid or amino acid sequence; (6) a polypeptide, such as a mIL-1Ra3 polypeptide, consisting of the amino acid sequence of amino acid residues 2 to 155, inclusive of Figure 9 (SEQ ID NO:16).

In a further aspect, the invention provides an isolated IL-11p polypeptide selected from the group consisting of: (1) a polypeptide comprising the amino acid sequence of amino acid residues 10 to 134, inclusive of Figure 5 (SEQ ID NO:10); (2) a polypeptide, such as an hIL-1Ra3 polypeptide, comprising the amino acid sequence of amino acid residues 2 to 155, inclusive of Figure 7 (SEQ ID NO: 13); and (3) a polypeptide, such as a mIL-1Ra3 polypeptide, comprising the amino acid sequence of amino acid residues from 2 to 155, inclusive of Figure 9 (SEQ ID NO:16).

In a still further aspect, the invention provides an isolated IL-1lp polypeptide that is the same as a mature polypeptide encoded by the cDNA insert of a vector selected from the group consisting of the vectors deposited as ATCC Deposit Nos. 203588, 203587, 203586, 203589, and 203590.

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In a still further aspect, the invention provides an isolated IL-1lp polypeptide that is the same as a mature polypeptide encoded by the cDNA insert of a vector selected from the group consisting of the vectors deposited as ATCC Deposit Nos. 203846, 203855 and 203973.

In a still further aspect, the invention provides an isolated polypeptide comprising the entire amino acid sequence encoded by the longest open reading frame in the cDNA insert of a vector selected from the group consisting of the vectors deposited as ATCC Deposit Nos. 203588, 203586, 203589 and 203590.

In a still further aspect, the invention provides an isolated polypeptide comprising the entire amino acid sequence encoded by the longest open reading frame in the cDNA insert of a vector selected from the group consisting of the vectors deposited as ATCC Deposit Nos. 203846, 203855 and 203973.

In another aspect, the invention provides a polypeptide that is produced by (i) hybridizing a test DNA molecule under stringent conditions with (a) a DNA molecule encoding an amino acid sequence selected from the group consisting of: (1) amino acid residues from at or about 37 to at or about 203, inclusive of Figure 2 (SEQ ID NO:5), (2) amino acid residues from at or about 15 to at or about 193, inclusive of Figure 3 (SEQ ID NO:7), (3) amino acid residues from at or about 2 to at or about 155, inclusive of Figure 7 (SEQ ID NO:13), and (4) amino acid residues from at or about 2 to at or about 155, inclusive of Figure 9 (SEQ ID NO:16), or (b) the complement of the DNA molecule of (a), and if the test DNA molecule encodes an IL-11p polypeptide that retains at least one biologic activity of a native sequence lL-1lp, such as the IL-1R binding activity of a native sequence hIL-1Ra3 or mIL-1Ra3, or the IL-18R binding activity of a native sequence hIL-1Ra1, hIL-1Ra1L, or hIL-1Ra1V, and if the test DNA molecule has at least at or about an 80% sequence identity, or at least at or about an 85% sequence identity, or at least at or about a 90% sequence identity, or at least at or about a 95% sequence identity to the DNA molecule of (a) or (b), (ii) culturing a host cell comprising the test DNA molecule under conditions suitable for expression of the IL-1lp polypeptide, and (iii) recovering the IL-1lp polypeptide from the cell culture.

In a still further aspect, the invention provides a polypeptide that is produced by (i) hybridizing a test DNA molecule under stringent conditions with (a) a DNA molecule encoding an amino acid sequence selected from the group consisting of: (1) amino acid residues from at or about 26 to at or about 207, inclusive of Figure 15 (SEQ ID NO:19), (2) amino acid residues from at or about 1 to at or about 218, inclusive of Figure 19 (SEQ ID NO:25), (3) amino acid residues from at or about 12 to at or about 218, inclusive of Figure 19 (SEQ ID NO:25), (4) amino acid residues from at or about 37 to at or about 218, inclusive of Figure 19 (SEQ ID NO:25), and (5) amino acid residues from at or about 46 to at or about 218, inclusive of Figure 19 (SEQ ID NO:25), and (5) amino acid residues from at or about 46 to at or about 218, inclusive of Figure 19 (SEQ ID NO:25), and inclusive of Figure 19 (SEQ ID NO:25), and (5) amino acid residues from at or about 46 to at or about 218, inclusive of Figure 19 (SEQ ID NO:25), and inclusive of Figure 19 (SEQ ID NO:25

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the test DNA molecule under conditions suitable for expression of the IL-1lp polypeptide, and (iii) recovering the IL-1lp polypeptide from the cell culture.

## A. Preparation of IL-11p

The description below relates primarily to production of IL-1lp by culturing cells transformed or transfected with a vector containing IL-1lp nucleic acid. It is, of course, contemplated that alternative methods, which are well known in the art, may be employed to prepare IL-1lp. For instance, the IL-1lp sequence, or portions thereof, may be produced by direct peptide synthesis using solid-phase techniques [see, e.g., Stewart et al., Solid-Phase Peptide Synthesis, W.H. Freeman Co., San Francisco, CA (1969); Merrifield, J. Am. Chem. Soc., 85:2149-2154 (1963)]. In vitro protein synthesis may be performed using manual techniques or by automation. Automated synthesis may be accomplished, for instance, using an Applied Biosystems Peptide Synthesizer (Foster City, CA) using manufacturer's instructions. Various portions of the IL-1lp may be chemically synthesized separately and combined using chemical or enzymatic methods to produce the full-length IL-1lp.

#### Isolation of DNA Encoding IL-1lp

DNA encoding IL-11p may be obtained from a cDNA library prepared from tissue believed to possess the IL-11p mRNA and to express it at a detectable level. Accordingly, human IL-11p DNA can be conveniently obtained from a cDNA library prepared from human tissue, such as described in the Examples. The IL-11p-encoding gene may also be obtained from a genomic library or by oligonucleotide synthesis.

Libraries can be screened with probes (such as antibodies to the IL-Ilp or oligonucleotides of at least about 20-80 bases) designed to identify the gene of interest or the protein encoded by it. Screening the cDNA or genomic library with the selected probe may be conducted using standard procedures, such as described in Sambrook et al., Molecular Cloning: A Laboratory Manual (New York: Cold Spring Harbor Laboratory Press, 1989). An alternative means to isolate the gene encoding IL-Ilp is to use PCR methodology [Sambrook et al., supra; Dieffenbach et al., PCR Primer: A Laboratory Manual (Cold Spring Harbor Laboratory Press, 1995)].

The Examples below describe techniques for screening a cDNA library. The oligonucleotide sequences selected as probes should be of sufficient length and sufficiently unambiguous that false positives are minimized. The oligonucleotide is preferably labeled such that it can be detected upon hybridization to DNA in the library being screened. Methods of labeling are well known in the art, and include the use of radiolabels like <sup>32</sup>P-labeled ATP, biotinylation or enzyme labeling. Hybridization conditions, including moderate stringency and high stringency, are provided in Sambrook et al., supra.

Sequences identified in such library screening methods can be compared and aligned to other known sequences deposited and available in public databases such as GenBank or other private sequence databases. Sequence identity (at either the amino acid or nucleotide level) within defined regions of the molecule or across the full-length sequence can be determined

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WO 00/39297 PCT/US99/30720

through sequence alignment using computer software programs such as BLAST, BLAST2, ALIGN-2, DNAstar, and INHERIT which employ various algorithms to measure homology.

Nucleic acid having protein coding sequence may be obtained by screening selected cDNA or genomic libraries using the deduced amino acid sequence disclosed herein for the first time, and, if necessary, using conventional primer extension procedures as described in Sambrook et al., supra, to detect precursors and processing intermediates of mRNA that may not have been reverse-transcribed into cDNA.

### 2. Selection and Transformation of Host Cells

Host cells are transfected or transformed with expression or cloning vectors described herein for IL-11p production and cultured in conventional nutrient media modified as appropriate for inducing promoters, selecting transformants, or amplifying the genes encoding the desired sequences. The culture conditions, such as media, temperature, pH and the like, can be selected by the skilled artisan without undue experimentation. In general, principles, protocols, and practical techniques for maximizing the productivity of cell cultures can be found in Mammalian Cell Biotechnology: A Practical Approach, M. Butler, ed. (IRL Press, 1991) and Sambrook et al., supra.

Methods of transfection are known to the ordinarily skilled artisan, for example, CaPO<sub>4</sub> and electroporation. Depending on the host cell used, transformation is performed using standard techniques appropriate to such cells. The calcium treatment employing calcium chloride, as described in Sambrook et al., supra, or electroporation is generally used for prokaryotes or other cells that contain substantial cell-wall barriers. Infection with Agrobacterium tumefaciens is used for transformation of certain plant cells, as described by Shaw et al., Gene, 23:315 (1983) and WO 89/05859 published 29 June 1989. For mammalian cells without such cell walls, the calcium phosphate precipitation method of Graham and van der Eb, Virology, 52:456-457 (1978) can be employed. General aspects of mammalian cell host system transformations have been described in U.S. Patent No. 4,399,216. Transformations into yeast are typically carried out according to the method of Van Solingen et al., J. Bact., 130:946 (1977) and Hsiao et al., Proc. Natl. Acad. Sci. (USA), 76:3829 (1979). However, other methods for introducing DNA into cells, such as by nuclear microinjection, electroporation, bacterial protoplast fusion with intact cells, or polycations, e.g., polybrene, polyornithine, may also be used. For various techniques for transforming mammalian cells, see Keown et al., Methods in Enzymology, 185:527-537 (1990) and Mansour et al., Nature, 336:348-352 (1988).

Suitable host cells for cloning or expressing the DNA in the vectors herein include prokaryote, yeast, or higher eukaryote cells. Suitable prokaryotes include but are not limited to eubacteria, such as Gram-negative or Gram-positive organisms, for example, Enterobacteriaceae such as E. coli. Various E. coli strains are publicly available, such as E. coli K12 strain MM294 (ATCC 31,446); E. coli X1776 (ATCC 31,537); E. coli strain W3110 (ATCC 27,325) and K5 772 (ATCC 53,635).

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In addition to prokaryotes, eukaryotic microbes such as filamentous fungi or yeast are suitable cloning or expression hosts for IL-11p-encoding vectors. Saccharomyces cerevisiae is a commonly used lower eukaryotic host microorganism.

Suitable host cells for the expression of glycosylated IL-1lp are derived from multicellular organisms. Examples of invertebrate cells include insect cells such as Drosophila S2 and Spodoptera Sf9, as well as plant cells. Examples of useful mammalian host cell lines include Chinese hamster ovary (CHO) and COS cells. More specific examples include monkey kidney CV1 line transformed by SV40 (COS-7, ATCC CRL 1651); human embryonic kidney line (293 cells subcloned for growth in suspension culture, Graham et al., J. Gen Virol., 36:59 (1977)); Chinese hamster ovary cells/-DHFR (CHO, Urlaub and Chasin, Proc. Natl. Acad. Sci. USA, 77:4216 (1980)); mouse sertoli cells (TM4, Mather, Biol. Reprod., 23:243-251 (1980)); human lung cells (W138, ATCC CCL 75); human liver cells (Hep G2, HB 8065); and mouse mammary tumor (MMT 060562, ATCC CCL51). The selection of the appropriate host cell is deemed to be within the skill in the art.

# Selection and Use of a Replicable Vector

The nucleic acid (e.g., cDNA or genomic DNA) encoding IL-11p may be inserted into a replicable vector for cloning (amplification of the DNA) or for expression. Various vectors are publicly available. The vector may, for example, be in the form of a plasmid, cosmid, viral particle, or phage. The appropriate nucleic acid sequence may be inserted into the vector by a variety of procedures. In general, DNA is inserted into an appropriate restriction endonuclease site(s) using techniques known in the art. Vector components generally include, but are not limited to, one or more of a signal sequence, an origin of replication, one or more marker genes, an enhancer element, a promoter, and a transcription termination sequence. Construction of suitable vectors containing one or more of these components employs standard ligation techniques which are known to the skilled artisan.

The IL-Ilp may be produced recombinantly not only directly, but also as a fusion polypeptide with a heterologous polypeptide, which may be a signal sequence or other polypeptide having a specific cleavage site at the N-terminus of the mature protein or polypeptide. In general, the signal sequence may be a component of the vector, or it may be a prokaryotic signal sequence may be the prokaryotic signal sequence selected, for example, from the group of the alkaline phosphatase, penicillinase, lpp, or heat-stable enterotoxin II leaders. For yeast secretion the signal sequence may be, e.g., the yeast invertase leader, alpha factor leader (including Saccharomyces and Kluyveromyces  $\alpha$ -factor leaders, the latter described in U.S. Patent No. 5,010,182), or acid phosphatase leader, the C. albicans glucoamylase leader (EP 362,179 published 4 April 1990), or the signal described in WO 90/13646 published 15 November 1990. In mammalian cell expression, mammalian signal sequences may be used to direct secretion of the protein, such as signal sequences from secreted polypeptides of the same or related species, as well as viral secretory leaders.

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WO 00/39297 PCT/US99/30720

Both expression and cloning vectors contain a nucleic acid sequence that enables the vector to replicate in one or more selected host cells. Such sequences are well known for a variety of bacteria, yeast, and viruses. The origin of replication from the plasmid pBR322 is suitable for most Gram-negative bacteria, the 2µ plasmid origin is suitable for yeast, and various viral origins (SV40, polyoma, adenovirus, VSV or BPV) are useful for cloning vectors in mammalian cells.

Expression and cloning vectors will typically contain a selection gene, also termed a selectable marker. Typical selection genes encode proteins that (a) confer resistance to antibiotics or other toxins, e.g., ampicillin, neomycin, methotrexate, or tetracycline, (b) complement auxotrophic deficiencies, or (c) supply critical nutrients not available from complex media, e.g., the gene encoding D-alanine racemase for Bacilli.

An example of suitable selectable markers for mammalian cells are those that enable the identification of cells competent to take up the IL-1lp-encoding nucleic acid, such as DHFR or thymidine kinase. An appropriate host cell when wild-type DHFR is employed is the CHO cell line deficient in DHFR activity, prepared and propagated as described by Urlaub et al., Proc. Natl. Acad. Sci. USA, 77:4216 (1980). A suitable selection gene for use in yeast is the trp1 gene present in the yeast plasmid YRp7 [Stinchcomb et al., Nature, 282:39 (1979); Kingsman et al., Gene, 7:141 (1979); Tschemper et al., Gene, 10:157 (1980)]. The trp1 gene provides a selection marker for a mutant strain of yeast lacking the ability to grow in tryptophan, for example, ATCC No. 44076 or PEP4-1 [Jones, Genetics, 85:12 (1977)].

Expression and cloning vectors usually contain a promoter operably linked to the IL-11p-encoding nucleic acid sequence to direct mRNA synthesis. Promoters recognized by a variety of potential host cells are well known. Promoters suitable for use with prokaryotic hosts include the β-lactamase and lactose promoter systems [Chang et al., Nature, 275:615 (1978); Goeddel et al., Nature, 281:544 (1979)], alkaline phosphatase, a tryptophan (trp) promoter system [Goeddel, Nucleic Acids Res., 8:4057 (1980); EP 36,776], and hybrid promoters such as the tac promoter deBoer et al., Proc. Natl. Acad. Sci. USA, 80:21-25 (1983)]. Promoters for use in bacterial systems also will contain a Shine-Dalgarno (S.D.) sequence operably linked to the DNA encoding IL-11b.

Examples of suitable promoting sequences for use with yeast hosts include the promoters for 3-phosphoglycerate kinase [Hitzeman et al., J. Biol. Chem., 255:2073 [1980]] or other glycolytic enzymes [Hess et al., J. Adv. Enzyme Reg., 7:149 [1968]; Holland, Biochemistry, 17:4900 (1978]], such as enolase, glyceraldehyde-3-phosphate dehydrogenase, hexokinase, pyruvate 'decarboxylase, phosphofructokinase, glucose-6-phosphate isomerase, 3-phosphoglycerate mutase, pyruvate kinase, triosephosphate isomerase, phosphoglucose isomerase, and glucokinase.

Other yeast promoters, which are inducible promoters having the additional advantage of transcription controlled by growth conditions, are the promoter regions for alcohol dehydrogenase 2, isocytochrome C, acid phosphatase, degradative enzymes associated with nitrogen metabolism, metallothionein, glyceraldehyde-3-phosphate dehydrogenase, and enzymes

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responsible for maltose and galactose utilization. Suitable vectors and promoters for use in yeast expression are further described in EP 73,657.

IL-11p transcription from vectors in mammalian host cells is controlled, for example, by promoters obtained from the genomes of viruses such as polyoma virus, fowlpox virus (UK 2,211,504 published 5 July 1989), adenovirus (such as Adenovirus 2), bovine papilloma virus, avian sarcoma virus, cytomegalovirus, a retrovirus, hepatitis-B virus and Simian Virus 40 (SV40), from heterologous mammalian promoters, e.g., the actin promoter or an immunoglobulin promoter, and from heat-shock promoters, provided such promoters are compatible with the host cell systems.

Transcription of a DNA encoding the IL-1lp by higher eukaryotes may be increased by inserting an enhancer sequence into the vector. Enhancers are cis-acting elements of DNA, usually about from 10 to 300 bp, that act on a promoter to increase its transcription. Many enhancer sequences are now known from mammalian genes (globin, elastase, albumin, α-fetoprotein, and insulin). Typically, however, one will use an enhancer from a eukaryotic cell virus. Examples include the SV40 enhancer on the late side of the replication origin (bp 100-270), the cytomegalovirus early promoter enhancer, the polyoma enhancer on the late side of the replication origin, and adenovirus enhancers. The enhancer may be spliced into the vector at a position 5' or 3' to the IL-1lp coding sequence, but is preferably located at a site 5' from the promoter.

Expression vectors used in eukaryotic host cells (yeast, fungi, insect, plant, animal, human, or nucleated cells from other multicellular organisms) will also contain sequences necessary for the termination of transcription and for stabilizing the mRNA. Such sequences are commonly available from the 5' and, occasionally 3', untranslated regions of eukaryotic or viral DNAs or cDNAs. These regions contain nucleotide segments transcribed as polyadenylated fragments in the untranslated portion of the mRNA encoding IL-11p.

Still other methods, vectors, and host cells suitable for adaptation to the synthesis of IL-11p in recombinant vertebrate cell culture are described in Gething et al., Nature, 293:620-625 (1981); Mantei et al., Nature, 281:40-46 (1979); EP 117,060; and EP 117,058.

## 4. Detecting Gene Amplification/Expression

Gene amplification and/or expression may be measured in a sample directly, for example, by conventional Southern blotting, Northern blotting to quantitate the transcription of mRNA [Thomas, Proc. Natl. Acad. Sci. USA, ZT:5201-5205 (1980)], dot blotting (DNA analysis), or in situ hybridization, using an appropriately labeled probe, based on the sequences provided herein. Alternatively, antibodies may be employed that can recognize specific duplexes, including DNA duplexes, RNA duplexes, and DNA-RNA hybrid duplexes or DNA-protein duplexes. The antibodies in turn may be labeled and the assay may be carried out where the duplex is bound to a surface, so that upon the formation of duplex on the surface, the presence of antibody bound to the duplex can be detected.

Gene expression, alternatively, may be measured by immunological methods, such as immunohistochemical staining of cells or tissue sections and assay of cell culture or body fluids,

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to quantitate directly the expression of gene product. Antibodies useful for immunohistochemical staining and/or assay of sample fluids may be either monoclonal or polyclonal, and may be prepared in any mammal. Conveniently, the antibodies may be prepared against a native sequence IL-11p polypeptide or against a synthetic peptide based on the DNA sequences provided herein or against exogenous sequence fused to IL-11p DNA and encoding a specific antibody epitope.

#### Purification of Polypeptide

Forms of IL-1lp may be recovered from culture medium or from host cell lysates. If membrane-bound, it can be released from the membrane using a suitable detergent solution (e.g. Triton-X 100) or by enzymatic cleavage. Cells employed in expression of IL-1lp can be disrupted by various physical or chemical means, such as freeze-thaw cycling, sonication, mechanical disruption, or cell lysing agents. It may be desired to purify IL-1lp from recombinant cell proteins or polypeptides. The following procedures are exemplary of suitable purification procedures: by fractionation on an ion-exchange column; ethanol precipitation; reverse phase HPLC; chromatography on silica or on a cation-exchange resin such as DEAE; chromatofocusing; SDS-PAGE; ammonium sulfate precipitation; gel filtration using, for example, Sephadex G-75; protein A Sepharose columns to remove contaminants such as IgG; and metal chelating columns to bind epitope-tagged forms of the IL-1lp. Various methods of protein purification may be employed and such methods are known in the art and described for example in Deutscher, Methods in Enzymology, 182 (1990); Scopes, Protein Purification: Principles and Practice, Springer-Verlag, New York (1982). The purification step(s) selected will depend, for example, on the nature of the production process used and the particular IL-1lp produced.

# B. Activity Assays for IL-11p Variants

The biological activity or activities of a particular IL-1lp variant polypeptide can be characterized using a variety of in vitro assays known in the art. For example, the ability of an hIL-1Ra3 variant polypeptide or a mIL-1Ra3 variant polypeptide to bind IL-1R can be assayed using a radioimmunoprecipitation assay wherein IL-1R extracellular domain (ECD) fused to the Fc region of human immunoglobulin G (IL-1R ECD-Fc) (which can be prepared, e.g., as described in Examples 9 and 10 below) is incubated in solution with radiolabeled hIL-1Ra3 variant polypeptide or mIL-1Ra3 variant polypeptide to form labeled complexes, followed by immunoprecipitation of the labeled complexes with goat anti-human IgG Fc and quantitation of radioactivity in the precipitate. In another example, an hIL-1Ra3 variant polypeptide-FLAG tag fusion protein-encoding DNA and an IL-1R ECD-Fc encoding DNA can be coexpressed in a host cell and secreted into the cell's culture medium, followed by immunoprecipitation of culture supernatant with protein G-sepharose and identification of bound hIL-1Ra3 variant polypeptide-FLAG tag fusion protein by immunoblotting with anti-FLAG monoclonal antibody, essentially as described in Example 9 below.

In another embodiment, the ability of an hIL-1Ra3 variant polypeptide or a mIL-1Ra3 variant polypeptide to inhibit the binding of IL-1 to IL-1R can be assayed using a competitive binding assay. For example, a radioimmunoprecipitation assay can be employed wherein IL-1R ECD-Fc is incubated in solution of radiolabeled IL-1 with or without unlabeled hIL-1Ra3 variant

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polypeptide or unlabeled mIL-1Ra3 variant polypeptide to form labeled or unlabeled complexes, followed by immunoprecipitation of complexes with anti-human IgG Fc and quantitation of radioactivity in the precipitate. If the presence of unlabeled hIL-1Ra3 variant polypeptide or unlabeled mIL-1Ra3 variant polypeptide in the incubation solution diminishes the radioactivity measured in the resulting immunoprecipitate, the hIL-1Ra3 variant polypeptide or mIL-1Ra3 variant polypeptide in question qualifies as an inhibitor of IL-1 binding to IL-1R. In yet another embodiment, IL-1R ECD-Fc and an hIL-1Ra3 variant-FLAG tag fusion protein or mIL-1Ra3 variant-FLAG tag fusion protein are obtained by recombinant expression in separate cell cultures (essentially as described in Example 10 below), IL-1 and IL-1R ECD-Fc are admixed together with or without the hIL-1Ra3 variant-FLAG tag fusion protein or mIL-1Ra3 variant-FLAG tag fusion protein and incubated in solution, the incubation solution is immunoprecipitated with protein G-sepharose, and the bound hlL-1Ra3 variant-FLAG tag fusion protein or mIL-1Ra3 variant-FLAG tag fusion protein is identified by immunoblotting with anti-FLAG monoclonal antibody. If the presence of IL-1 in the incubation solution diminishes the signal detected by anti-FLAG immunoblotting, the hlL-1Ra3 variant polypeptide or mIL-1Ra3 variant polypeptide in question qualifies as an inhibitor of IL-1 binding to IL-1R.

Similarly, the biological activity or activities of a particular hIL-1Ra1 variant polypeptide can be determined by using a variety of in vitro assays known in the art. For example, the ability of an hIL-1Ra1 variant polypeptide to bind IL-18R can be assayed using a radioimmunoprecipitation assay wherein IL-18R extracellular domain (ECD) fused to the Foregion of human immunoglobulin G (IL-18R ECD-Fc) (which can be prepared, e.g., as described in Examples 9 and 10 below) is incubated in solution with radiolabeled hIL-1Ra1 variant polypeptide to form labeled complex, followed by immunoprecipitation of the labeled complex with goat anti-human lgG Fc and quantitation of radioactivity in the precipitate. In another example, an hIL-1Ra1 variant polypeptide-FLAG tag fusion protein-encoding DNA and an IL-18R ECD-Fc encoding DNA can be coexpressed in a host cell and secreted into the cell's culture medium, followed by immunoprecipitation of culture supermatant with protein G-sepharose and identification of bound hIL-1Ra1 variant polypeptide-FLAG tag fusion protein by immunoblotting with anti-FLAG monoclonal antibody, essentially as described in Example 9 below.

In another embodiment, the ability of an hIL-1Ra1 variant polypeptide to inhibit the binding of IL-18 to IL-18R can be assayed using a competitive binding assay. For example, a radioimmunoprecipitation assay can be employed wherein IL-18R ECD-Fc is incubated in solution of radiolabeled IL-18 with or without unlabeled hIL-1Ra1 variant polypeptide to form labeled or unlabeled complexes, followed by immunoprecipitation of complexes with anti-human lgG Fc and quantitation of radioactivity in the precipitate. If the presence of unlabeled hIL-1Ra1 variant polypeptide in the incubation solution diminishes the radioactivity measured in the resulting immunoprecipitate, the hIL-1Ra1 variant polypeptide in question qualifies as an inhibitor of IL-18 binding to IL-18R. In yet another embodiment, IL-18R ECD-Fc and an hIL-1Ra1 variant-FLAG tag fusion protein are obtained by recombinant expression in separate cell cultures (essentially as described in Example 10 below), IL-18 and IL-18R ECD-Fc are admixed together with or without the hIL-1Ra1 variant-FLAG tag fusion protein and incubated in

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solution, the incubation solution is immunoprecipitated with protein G-sepharose, and the bound hIL-1Ra1 variant-FLAG tag fusion protein is identified by immunoblotting with anti-FLAG monoclonal antibody. If the presence of IL-18 in the incubation solution diminishes the signal detected by anti-FLAG immunoblotting, the hIL-1Ra1 variant polypeptide in question qualifies as an inhibitor of IL-18 binding to IL-18R.

#### C. Uses for IL-11p

Nucleotide sequences (or their complement) encoding IL-1lp have various applications in the art of molecular biology, including uses as hybridization probes, in chromosome and gene mapping and in the generation of anti-sense RNA and DNA. IL-1lp nucleic acid will also be useful for the preparation of IL-1lp polypeptides by the recombinant techniques described herein.

The full-length native sequence IL-11p genes of Figure 1 (SEQ ID NO:1), Figure 2 (SEQ ID NO:4), Figure 3 (SEQ ID NO:6), Figure 5 (SEQ ID NO:9), Figure 7 (SEQ ID NO:12), Figure 9 (SEQ ID NO:15), Figure 15 (SEQ ID NO:18), and Figure 16 (SEQ ID NO:20), and Figure 19 (SEQ ID NO:24), or portions thereof, may be used as hybridization probes for a cDNA library to isolate the full-length IL-1lp gene or to isolate still other genes (for instance, those encoding naturallyoccurring variants of IL-1lp or IL-1lp from other species) which have a desired sequence identity to the IL-1lp sequence disclosed in Figure 1 (SEQ ID NO:1), Figure 2 (SEQ ID NO:4), Figure 3 (SEQ ID NO:6), Figure 5 (SEQ ID NO:9), Figure 7 (SEQ ID NO:12), Figure 9 (SEQ ID NO:15), Figure 15 (SEQ ID NO:18), Figure 16 (SEQ ID NO:20), or Figure 19 (SEQ ID NO:24). Optionally, the length of the probes will be about 20 to about 50 bases. The hybridization probes may be derived from the nucleotide sequence of Figure 1 (SEQ ID NO:1), Figure 2 (SEQ ID NO:4), Figure 3 (SEQ ID NO:6), Figure 5 (SEQ ID NO:9), Figure 7 (SEQ ID NO:12), Figure 9 (SEQ ID NO:15), Figure 15 (SEQ ID NO:18), Figure 16 (SEQ ID NO:20) or Figure 19 (SEQ ID NO:24), or from genomic sequences including promoters, enhancer elements and introns of native sequence IL-1lp. By way of example, a screening method will comprise isolating the coding region of the IL-1lp gene using the known DNA sequence to synthesize a selected probe of about 40 bases. Hybridization probes may be labeled by a variety of labels, including radionucleotides such as 32P or 35S, or enzymatic labels such as alkaline phosphatase coupled to the probe via avidin/biotin coupling systems. Labeled probes having a sequence complementary to that of the IL-11p gene of the present invention can be used to screen libraries of human cDNA, genomic DNA or mRNA to determine which members of such libraries the probe hybridizes to. Hybridization techniques are described in further detail in the Examples below.

The probes may also be employed in PCR techniques to generate a pool of sequences for identification of closely related IL-1lp coding sequences.

Nucleotide sequences encoding an IL-IIp can also be used to construct hybridization probes for mapping the gene which encodes that IL-IIp and for the genetic analysis of individuals with genetic disorders. The nucleotide sequences provided herein may be mapped to a chromosome and specific regions of a chromosome using known techniques, such as in situ hybridization, linkage analysis against known chromosomal markers, and hybridization screening with libraries.

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When the coding sequences for IL-1lp encode a protein which binds to another protein (example, where the IL-1lp binds to an IL-1 receptor or IL-18 receptor), the IL-1lp can be used in assays to identify the other proteins or molecules involved in the binding interaction. By such methods, inhibitors of the receptor/ligand binding interaction can be identified. Proteins involved in such binding interactions can also be used to screen for peptide or small molecule inhibitors or agonists of the binding interaction. Screening assays can be designed to find lead compounds that mimic the biological activity of a native IL-1lp or a receptor for IL-1lp. Such screening assays will include assays amenable to high-throughput screening of chemical libraries, making them particularly suitable for identifying small molecule drug candidates. Small molecules contemplated include synthetic organic or inorganic compounds. The assays can be performed in a variety of formats, including protein-protein binding assays, biochemical screening assays, immunoassays and cell based assays, which are well characterized in the art.

Nucleic acids which encode IL-11p or its modified forms can also be used to generate either transgenic animals or "knock out" animals which, in turn, are useful in the development and screening of therapeutically useful reagents. A transgenic animal (e.g., a mouse or rat) is an animal having cells that contain a transgene, which transgene was introduced into the animal or an ancestor of the animal at a prenatal, e.g., an embryonic stage. A transgene is a DNA which is integrated into the genome of a cell from which a transgenic animal develops. In one embodiment, cDNA encoding IL-11p can be used to clone genomic DNA encoding IL-11p in accordance with established techniques and the genomic sequences used to generate transgenic animals that contain cells which express DNA encoding IL-11p. Methods for generating transgenic animals, particularly animals such as mice or rats, have become conventional in the art and are described, for example, in U.S. Patent Nos. 4,736,866 and 4,870,009. Typically, particular cells would be targeted for IL-1lp transgene incorporation with tissue-specific enhancers. Transgenic animals that include a copy of a transgene encoding IL-1lp introduced into the germ line of the animal at an embryonic stage can be used to examine the effect of increased expression of DNA encoding IL-11p. Such animals can be used as tester animals for reagents thought to confer protection from, for example, pathological conditions associated with its overexpression. In accordance with this facet of the invention, an animal is treated with the reagent and a reduced incidence of the pathological condition, compared to untreated animals bearing the transgene, would indicate a potential therapeutic intervention for the pathological condition.

Alternatively, non-human homologues of IL-1lp can be used to construct an IL-1lp "knock out" animal which has a defective or altered gene encoding IL-1lp as a result of homologous recombination between the endogenous gene encoding IL-1lp and altered genomic DNA encoding IL-1lp introduced into an embryonic cell of the animal. For example, cDNA encoding IL-1lp can be used to clone genomic DNA encoding IL-1lp in accordance DNA encoding IL-1lp can be deleted or replaced with another gene, such as a gene encoding a selectable marker which can be used to monitor integration. Typically, several kilobases of unaltered flanking DNA (both at the 5' and 3' ends) are included in the vector [see e.g., Thomas and Capecchi, Cell, 51:503 (1987) for a description

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of homologous recombination vectors]. The vector is introduced into an embryonic stem cell line (e.g., by electroporation) and cells in which the introduced DNA has homologously recombined with the endogenous DNA are selected [see e.g., Li et al., Cell, 69:915 (1992)]. The selected cells are then injected into a blastocyst of an animal (e.g., a mouse or rat) to form aggregation chimeras [see e.g., Bradley, in Teratocarcinomas and Embryonic Stem Cells: A Practical Approach, E. J. Robertson, ed. (IRL, Oxford, 1987), pp. 113-152]. A chimeric embryo can then be implanted into a suitable pseudopregnant female foster animal and the embryo brought to term to create a "knock out" animal. Progeny harboring the homologously recombined DNA in their germ cells can be identified by standard techniques and used to breed animals in which all cells of the animal contain the homologously recombined DNA. Knockout animals can be characterized for instance, for their ability to defend against certain pathological conditions and for their development of pathological conditions due to absence of the IL-1lp polypeptide.

Nucleic acid encoding the IL-11p polypeptides may also be used in gene therapy. In gene therapy applications, genes are introduced into cells in order to achieve in vivo synthesis of a therapeutically effective genetic product, for example for replacement of a defective gene. "Gene therapy" includes both conventional gene therapy where a lasting effect is achieved by a single treatment, and the administration of gene therapeutic agents, which involves the one time or repeated administration of a therapeutically effective DNA or mRNA. Antisense RNAs and DNAs can be used as therapeutic agents for blocking the expression of certain genes in vivo. It has already been shown that short antisense oligonucleotides can be imported into cells where they act as inhibitors, despite their low intracellular concentrations caused by their restricted uptake by the cell membrane. (Zamecnik et al., Proc. Natl. Acad. Sci. USA §3, 4143-4146 [1986]). The oligonucleotides can be modified to enhance their uptake, e.g. by substituting their negatively charged phosphodiester groups by uncharged groups.

There are a variety of techniques available for introducing nucleic acids into viable cells. The techniques vary depending upon whether the nucleic acid is transferred into cultured cells in vitro, or in vivo in the cells of the intended host. Techniques suitable for the transfer of nucleic acid into mammalian cells in vitro include the use of liposomes, electroporation, microinjection, cell fusion, DEAE-dextran, the calcium phosphate precipitation method, etc. The currently preferred in vivo gene transfer techniques include transfection with viral (typically retroviral) vectors and viral coat protein-liposome mediated transfection (Dzau et al., Trends in Biotechnology 11, 205-210 [1993]). In some situations it is desirable to provide the nucleic acid source with an agent that targets the target cells, such as an antibody specific for a cell surface membrane protein or the target cell, a ligand for a receptor on the target cell, etc. Where liposomes are employed, proteins which bind to a cell surface membrane protein associated with endocytosis may be used for targeting and/or to facilitate uptake, e.g. capsid proteins or fragments thereof tropic for a particular cell type, antibodies for proteins which undergo internalization in cycling, proteins that target intracellular localization and enhance intracellular half-life. The technique of receptor-mediated endocytosis is described, for example, by Wu et al., J. Biol. Chem. 262, 4429-4432 (1987); and Wagner et al., Proc. Natl. Acad. Sci. USA 87, 3410-

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WO 00/39297 PCT/US99/30720

3414 (1990). For review of gene marking and gene therapy protocols see Anderson *et al.*, *Science* 256, 808-813 (1992).

The IL-1lp polypeptides of the present invention can be formulated according to known methods to prepare pharmaceutically useful compositions, whereby the IL-1lp product hereof is combined in admixture with a pharmaceutically acceptable carrier vehicle. Therapeutic formulations are prepared for storage by mixing the active ingredient having the desired degree of purity with optional physiologically acceptable carriers, excipients or stabilizers (Remington's Pharmaceutical Sciences 16th edition, Osol, A. Ed. (1980)), in the form of lyophilized formulations or aqueous solutions. Acceptable carriers, excipients or stabilizers are nontoxic to recipients at the dosages and concentrations employed, and include buffers such as phosphate, citrate and other organic acids; antioxidants including ascorbic acid; low molecular weight (less than about 10 residues) polypeptides; proteins, such as serum albumin, gelatin or immunoglobulins; hydrophilic polymers such as polyvinylpyrrolidone, amino acids such as glycine, glutamine, asparagine, arginine or lysine; monosaccharides, disaccharides and other carbohydrates including glucose, mannose, or dextrins; chelating agents such as EDTA; sugar alcohols such as mannitol or sorbitol; salt-forming counterions such as sodium; and/or nonionic surfactants such as Tween, Pluronics or PEG.

The formulations to be used for *in vivo* administration must be sterile. This is readily accomplished by filtration through sterile filtration membranes, prior to or following lyophilization and reconstitution.

Therapeutic compositions herein generally are placed into a container having a sterile access port, for example, an intravenous solution bag or vial having a stopper pierceable by a hypodermic injection needle.

The route of administration is in accord with known methods, e.g. injection or infusion by intravenous, intraperitoneal, intracerebral, intramuscular, intraocular, intraarterial or intralesional routes, topical administration, or by sustained release systems.

Dosages and desired drug concentrations of pharmaceutical compositions of the present invention may vary depending on the particular use envisioned. The determination of the appropriate dosage or route of administration is well within the skill of an ordinary physician. Animal experiments provide reliable guidance for the determination of effective doses for human therapy. Interspecies scaling of effective doses can be performed following the principles laid down by Mordenti, J. and Chappell, W. "The use of interspecies scaling in toxicokinetics' In Toxicokinetics and New Drug Development, Yacobi et al., Eds., Pergamon Press, New York 1989, pp. 42-96.

An "effective amount" of the IL-1lp to be employed therapeutically will depend, for example, upon the therapeutic objectives, the route of administration, and the condition of the patient. Accordingly, it will be necessary for the therapeut to titer the dosage and modify the route of administration as required to obtain the optimal therapeutic effect. Typically, the clinician will administer the IL-1lp until a dosage is reached that achieves the desired effect. The progress of this therapy is easily monitored by conventional assays.

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In one embodiment, the invention provides a method for treating an IL-1-mediated disorder comprising administering to a mammal, such as human, in need of such treatment an effective amount of an IL-11p, such as a native sequence IL-11p.

In another embodiment, the invention provides a method for treating an IL-1-mediated disorder comprising administering to a mammal, such as human, in need of such treatment an effective amount of an IL-1lp selected from the group consisting of hIL-1Ra1, hIL-1Ra1L, hIL-1Ra1S, hIL-1Ra2, hIL-1Ra3, and mIL-1Ra3.

In another embodiment, the invention provides a method for treating an IL-1-mediated disorder comprising administering to a mammal, such as human, in need of such treatment an effective amount of an hIL-11p, such as a native sequence hIL-11p, e.g. native sequence hIL-11a3.

In another embodiment, the invention provides a method for treating an IL-18-mediated disorder comprising administering to a mammal, such as human, in need of such treatment an effective amount of an IL-11p, such as a native sequence IL-11p.

In another embodiment, the invention provides a method for treating an IL-18-mediated disorder comprising administering to a mammal, such as human, in need of such treatment an effective amount of an IL-1lp selected from the group consisting of hIL-1Ra1, hIL-1Ra1L, hIL-1Ra1V, hIL-1Ra1S, hIL-1Ra2, hIL-1Ra3, and mIL-1Ra3.

In another embodiment, the invention provides a method for treating an IL-18-mediated disorder comprising administering to a mammal, such as human, in need of such treatment an effective amount of an hIL-1lp, such as a native sequence hIL-1lp, e.g. native sequence hIL-1Ra1.

In another embodiment, the invention provides a method for treating an IL-18-mediated disorder comprising administering to a mammal, such as human, in need of such treatment an effective amount of an hIL-1Ra1L, such as a native sequence hIL-1Ra1L, or an effective amount of an hIL-1Ra1V, such as a native sequence hIL-1Ra1V, or an effective amount of an hIL-1Ra1S, such as a native sequence hIL-1Ra1S.

In one embodiment, the invention provides a method for treating an inflammatory disorder comprising administering to a mammal, such as human, in need of such treatment an effective amount of an IL-11p, such as a native sequence IL-11p.

In another embodiment, the invention provides a method for treating an inflammatory disorder comprising administering to a mammal, such as human, in need of such treatment an effective amount of an IL-1lp selected from the group consisting of hIL-1Ra1, hIL-1Ra1L, hIL-1Ra1V, hIL-1Ra1S, hIL-1Ra2, hIL-1Ra3, and mIL-1Ra3.

In another embodiment, the invention provides a method for treating an inflammatory disorder comprising administering to a mammal, such as human, in need of such treatment an effective amount of an hIL-1lp, such as a native sequence hIL-1lp, e.g. native sequence hIL-1Ra1 or hIL-1Ra3.

In another embodiment, the invention provides a method for treating an inflammatory disorder comprising administering to a mammal, such as human, in need of such treatment an effective amount of an hIL-1Ra1L, such as a native sequence hIL-1Ra1L, or an effective amount of an hIL-1Ra1V, such as a native sequence hIL-1Ra1V, or an effective amount of an hIL-1Ra1S, such as a native sequence hIL-1Ra1S.

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WO 90/39297 PCT/US99/30720

In another embodiment, the invention provides a method for treating asthma comprising administering to a mammal, such as human, in need of such treatment an effective amount of an IL-1lp, such as a native sequence IL-1lp.

In another embodiment, the invention provides a method for treating asthma comprising administering to a mammal, such as human, in need of such treatment an effective amount of an IL-1lp selected from the group consisting of hIL-1Ra1, hIL-1Ra1L, hIL-1Ra1V, hIL-1Ra1S, hIL-1Ra2, hIL-1Ra3, and mIL-1Ra3.

In another embodiment, the invention provides a method for treating asthma comprising administering to a mammal, such as human, in need of such treatment an effective amount of an hIL-1lp, such as a native sequence hIL-1lp, e.g. native sequence hIL-1Ra1 or hIL-1Ra3.

In another embodiment, the invention provides a method for treating asthma comprising administering to a mammal, such as human, in need of such treatment an effective amount of an hlL-1Ra1L, such as a native sequence hlL-1Ra1L, or an effective amount of an hlL-1Ra1V, such as a native sequence hlL-1Ra1V, or an effective amount of an hlL-1Ra1S, such as a native sequence hlL-1Ra1S.

In another embodiment, the invention provides a method for treating rheumatoid arthritis comprising administering to a mammal, such as human, in need of such treatment an effective amount of an IL-1lp, such as a native sequence IL-1lp.

In another embodiment, the invention provides a method for treating rheumatoid arthritis comprising administering to a mammal, such as human, in need of such treatment an effective amount of an IL-1lp selected from the group consisting of hIL-1Ra1, hIL-1Ra1L, hIL-1Ra1V, hIL-1Ra1S, hIL-1Ra2, hIL-1Ra3, and mIL-1Ra3.

In another embodiment, the invention provides a method for treating rheumatoid arthritis comprising administering to a mammal, such as human, in need of such treatment an effective amount of an hIL-1lp, such as a native sequence hIL-1lp, e.g. native sequence hIL-1Ra1 or hIL-1Ra3.

In another embodiment, the invention provides a method for treating rheumatoid arthritis comprising administering to a mammal, such as human, in need of such treatment an effective amount of an hIL-1Ra1L, such as a native sequence hIL-1Ra1L, or an effective amount of an hIL-1Ra1V, such as a native sequence hIL-1Ra1V, or an effective amount of an hIL-1Ra1S, such as a native sequence hIL-1Ra1S.

In another embodiment, the invention provides a method for treating osteoarthritis comprising administering to a mammal, such as human, in need of such treatment an effective amount of an IL-11p, such as a native sequence IL-11p.

In another embodiment, the invention provides a method for treating osteoarthritis comprising administering to a mammal, such as human, in need of such treatment an effective amount of an IL-1lp selected from the group consisting of hIL-1Ra1, hIL-1Ra1L, hIL-1Ra1V, hIL-1Ra1S, hIL-1Ra2, hIL-1Ra3, and mIL-1Ra3.

In another embodiment, the invention provides a method for treating osteoarthritis comprising administering to a mammal, such as human, in need of such treatment an effective

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amount of an hiL-1lp, such as a native sequence hIL-1lp, e.g. native sequence hIL-1Ra1 or hIL-

In another embodiment, the invention provides a method for treating osteoarthritis comprising administering to a mammal, such as human, in need of such treatment an effective amount of an hIL-1Ra1L, such as a native sequence hIL-1Ra1L, or an effective amount of an hIL-1Ra1V, such as a native sequence hIL-1Ra1V, or an effective amount of an hIL-1Ra1S, such as a native sequence hIL-1Ra1S.

In another embodiment, the invention provides a method for treating sepsis comprising administering to a mammal, such as human, in need of such treatment an effective amount of an IL-11p, such as a native sequence IL-11p.

In another embodiment, the invention provides a method for treating sepsis comprising administering to a mammal, such as human, in need of such treatment an effective amount of an IL-1lp selected from the group consisting of hIL-1Ra1, hIL-1Ra1L, hIL-1Ra1V, hIL-1Ra1S, hIL-1Ra2, hIL-1Ra3, and mIL-1Ra3.

In another embodiment, the invention provides a method for treating sepsis comprising administering to a mammal, such as human, in need of such treatment an effective amount of an hIL-11p, such as a native sequence hIL-11p, e.g. native sequence hIL-1Ra1 or hIL-1Ra3.

In another embodiment, the invention provides a method for treating sepsis comprising administering to a mammal, such as human, in need of such treatment an effective amount of an hIL-1Ra1L, such as a native sequence hIL-1Ra1L, or an effective amount of an hIL-1Ra1V, such as a native sequence hIL-1Ra1V, or an effective amount of an hIL-1Ra1S, such as a native sequence hIL-1Ra1S.

In another embodiment, the invention provides a method for treating acute lung injury comprising administering to a mammal, such as human, in need of such treatment an effective amount of an IL-11p, such as a native sequence IL-11p.

In another embodiment, the invention provides a method for treating acute lung injury comprising administering to a mammal, such as human, in need of such treatment an effective amount of an IL-11p selected from the group consisting of hIL-1Ra1, hIL-1Ra1L, hIL-1Ra1V, hIL-1Ra1S, hIL-1Ra2, hIL-1Ra3, and mIL-1Ra3.

In another embodiment, the invention provides a method for treating acute lung injury comprising administering to a mammal, such as human, in need of such treatment an effective amount of an hIL-1lp, such as a native sequence hIL-1lp, e.g. native sequence hIL-1Ra1 or hIL-1Ra3.

In another embodiment, the invention provides a method for treating acute lung injury comprising administering to a mammal, such as human, in need of such treatment an effective amount of an hIL-1Ra1L, such as a native sequence hIL-1Ra1L, or an effective amount of an hIL-1Ra1V, such as a native sequence hIL-1Ra1V, or an effective amount of an hIL-1Ra1S, such as a native sequence hIL-1Ra1S.

In another embodiment, the invention provides a method for treating adult respiratory distress syndrome comprising administering to a mammal, such as human, in need of such treatment an effective amount of an IL-1lp, such as a native sequence IL-1lp.

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In another embodiment, the invention provides a method for treating adult respiratory distress syndrome comprising administering to a mammal, such as human, in need of such treatment an effective amount of an IL-11p selected from the group consisting of hIL-1Ra1, hIL-1Ra1L, hIL-1Ra1V, hIL-1Ra1S, hIL-1Ra2, hIL-1Ra3, and mIL-1Ra3.

In another embodiment, the invention provides a method for treating adult respiratory distress syndrome comprising administering to a mammal, such as human, in need of such treatment an effective amount of an hIL-1lp, such as a native sequence hIL-1lp, e.g. native sequence hIL-1Ra1 or hIL-1Ra3.

In another embodiment, the invention provides a method for treating adult respiratory distress syndrome comprising administering to a mammal, such as human, in need of such treatment an effective amount of an hlL-1Ra1L, such as a native sequence hlL-1Ra1L, or an effective amount of an hlL-1Ra1V, such as a native sequence hlL-1Ra1V, or an effective amount of an hlL-1Ra1S, such as a native sequence hlL-1Ra1S.

In another embodiment, the invention provides a method for treating idiopathic pulmonary fibrosis comprising administering to a mammal, such as human, in need of such treatment an effective amount of an IL-1lp, such as a native sequence IL-1lp.

In another embodiment, the invention provides a method for treating idiopathic pulmonary fibrosis comprising administering to a mammal, such as human, in need of such treatment an effective amount of an IL-1lp selected from the group consisting of hIL-1Ra1, hIL-1Ra1L, hIL-1Ra1V, hIL-1Ra1S, hIL-1Ra2, hIL-1Ra3, and mIL-1Ra3.

In another embodiment, the invention provides a method for treating idiopathic pulmonary fibrosis comprising administering to a mammal, such as human, in need of such treatment an effective amount of an hIL-1lp, such as a native sequence hIL-1lp, e.g. native sequence hIL-1Ra1 or hIL-1Ra3.

In another embodiment, the invention provides a method for treating idiopathic pulmonary fibrosis comprising administering to a mammal, such as human, in need of such treatment an effective amount of an hIL-1Ra1L, such as a native sequence hIL-1Ra1L, or an effective amount of an hIL-1Ra1V, such as a native sequence hIL-1Ra1V, or an effective amount of an hIL-1Ra1S. such as a native sequence hIL-1Ra1S.

In another embodiment, the invention provides a method for treating an ischemic reperfusion disease, such as surgical tissue reperfusion injury, stroke, myocardial ischemia, or acute myocardial infarction, comprising administering to a mammal, such as human, in need of such treatment an effective amount of an IL-1lp, such as a native sequence IL-1lp.

In another embodiment, the invention provides a method for treating an ischemic reperfusion disease, such as surgical tissue reperfusion injury, stroke, myocardial ischemia, or acute myocardial infarction, comprising administering to a mammal, such as human, in need of such treatment an effective amount of an IL-Ilp selected from the group consisting of hIL-IRa1, hIL-IRa1L, hIL-IRa1V, hIL-IRa1S, hIL-IRa2, hIL-IRa3, and mIL-IRa3.

In another embodiment, the invention provides a method for treating an ischemic reperfusion disease, such as surgical tissue reperfusion injury, stroke, myocardial ischemia, or acute myocardial infarction, comprising administering to a mammal, such as human, in need

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of such treatment an effective amount of an hIL-1lp, such as a native sequence hIL-1lp, e.g. native sequence hIL-1Ra1 or hIL-1Ra3.

In another embodiment, the invention provides a method for treating an ischemic reperfusion disease, such as surgical tissue reperfusion injury, stroke, myocardial ischemia, or acute myocardial infarction, comprising administering to a mammal, such as human, in need of such treatment an effective amount of an hIL-1Ra1L, such as a native sequence hIL-1Ra1L, or an effective amount of an hIL-1Ra1V, such as a native sequence hIL-1Ra1V, or an effective amount of an hIL-1Ra1S, such as a native sequence hIL-1Ra1S.

In another embodiment, the invention provides a method for treating psoriasis comprising administering to a mammal, such as human, in need of such treatment an effective amount of an IL-1lp, such as a native sequence IL-1lp.

In another embodiment, the invention provides a method for treating psoriasis comprising administering to a mammal, such as human, in need of such treatment an effective amount of an IL-1lp selected from the group consisting of hIL-1Ra1, hIL-1Ra1L, hIL-1Ra1V, hIL-1Ra1S, hIL-1Ra2, hIL-1Ra3, and mIL-1Ra3.

In another embodiment, the invention provides a method for treating psoriasis comprising administering to a mammal, such as human, in need of such treatment an effective amount of an hIL-1lp, such as a native sequence hIL-1lp, e.g. native sequence hIL-1Ra1 or hIL-1Ra3.

In another embodiment, the invention provides a method for treating psoriasis comprising administering to a mammal, such as human, in need of such treatment an effective amount of an hIL-1Ra1L, such as a native sequence hIL-1Ra1L, or an effective amount of an hIL-1Ra1S, such as a native sequence hIL-1Ra1V, or an effective amount of an hIL-1Ra1S, such as a native sequence hIL-1Ra1S.

In another embodiment, the invention provides a method for treating graft-versus-host disease (GVHD) comprising administering to a mammal, such as human, in need of such treatment an effective amount of an IL-1lp, such as a native sequence IL-1lp.

In another embodiment, the invention provides a method for treating graft-versus-host disease (GVHD) comprising administering to a mammal, such as human, in need of such treatment an effective amount of an IL-1lp selected from the group consisting of hIL-1Ra1, hIL-1Ra1L, hIL-1Ra1V, hIL-1Ra1S, hIL-1Ra2, hIL-1Ra3, and mIL-1Ra3.

In another embodiment, the invention provides a method for treating graft-versus-host disease (GVHD) comprising administering to a mammal, such as human, in need of such treatment an effective amount of an hIL-1lp, such as a native sequence hIL-1lp, e.g. native sequence hIL-1Ra1 or hIL-1Ra3.

In another embodiment, the invention provides a method for treating graft-versus-host disease (GVHD) comprising administering to a mammal, such as human, in need of such treatment an effective amount of an hIL-1Ra1L, such as a native sequence hIL-1Ra1L, or an effective amount of an hIL-1Ra1V, such as a native sequence hIL-1Ra1V, or an effective amount of an hIL-1Ra1S, such as a native sequence hIL-1Ra1S.

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In another embodiment, the invention provides a method for treating an inflammatory bowel disease such as ulcerative colitis, comprising administering to a mammal, such as human, in need of such treatment an effective amount of an IL-1lp, such as a native sequence IL-1lp.

In another embodiment, the invention provides a method for treating an inflammatory bowel disease such as ulcerative colitis, comprising administering to a mammal, such as human, in need of such treatment an effective amount of an IL-1lp selected from the group consisting of hIL-1Ra1, hIL-1Ra1L, hIL-1Ra1V, hIL-1Ra1S, hIL-1Ra3, and mIL-1Ra3.

In another embodiment, the invention provides a method for treating an inflammatory bowel disease such as ulcerative colitis, comprising administering to a mammal, such as human, in need of such treatment an effective amount of an hIL-1lp, such as a native sequence hIL-1lp, e.g. native sequence hIL-1Ra1 or hIL-1Ra3.

In another embodiment, the invention provides a method for treating an inflammatory bowel disease such as ulcerative colitis, comprising administering to a mammal, such as human, in need of such treatment an effective amount of an hIL-1Ra1L, such as a native sequence hIL-1Ra1L, or an effective amount of an hIL-1Ra1V, such as a native sequence hIL-1Ra1V, or an effective amount of an hIL-1Ra1S, such as a native sequence hIL-1Ra1S.

## D. <u>Anti-IL-1lp Antibodies</u>

The present invention further provides anti-IL-1lp antibodies. Exemplary antibodies include polyclonal, monoclonal, humanized, bispecific, and heteroconjugate antibodies.

### Polyclonal Antibodies

The anti-IL-11p antibodies may comprise polyclonal antibodies. Methods of preparing polyclonal antibodies are known to the skilled artisan. Polyclonal antibodies can be raised in a mammal, for example, by one or more injections of an immunizing agent and, if desired, an adjuvant. Typically, the immunizing agent and/or adjuvant will be injected in the mammal by multiple subcutaneous or intraperitoneal injections. The immunizing agent may include the IL-11p polypeptide or a fusion protein thereof. It may be useful to conjugate the immunizing agent to a protein known to be immunogenic in the mammal being immunized. Examples of such immunogenic proteins include but are not limited to keyhole limpet hemocyanin, serum albumin, bovine thyroglobulin, and soybean trypsin inhibitor. Examples of adjuvants which may be employed include Freund's complete adjuvant and MPL-TDM adjuvant (monophosphoryl Lipid A, synthetic trehalose dicorynomycolate). The immunization protocol may be selected by one skilled in the art without undue experimentation.

#### Monoclonal Antibodies

The anti-IL-11p antibodies may, alternatively, be monoclonal antibodies. Monoclonal antibodies may be prepared using hybridoma methods, such as those described by Kohler and Milstein, Nature, 256:495 (1975). In a hybridoma method, a mouse, hamster, or other appropriate host animal, is typically immunized with an immunizing agent to elicit lymphocytes that produce or are capable of producing antibodies that will specifically bind to the immunizing agent. Alternatively, the lymphocytes may be immunized in vitro.

The immunizing agent will typically include the IL-1lp polypeptide or a fusion protein thereof. Generally, either peripheral blood lymphocytes ("PBLs") are used if cells of human origin

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WO 00/39297 PCT/US99/30720

are desired, or spleen cells or lymph node cells are used if non-human mammalian sources are desired. The lymphocytes are then fused with an immortalized cell line using a suitable fusing agent, such as polyethylene glycol, to form a hybridoma cell [Goding, Monoclonal Antibodies: Principles and Practice, Academic Press, (1986) pp. 59-103]. Immortalized cell lines are usually transformed mammalian cells, particularly myeloma cells of rodent, bovine and human origin. Usually, rat or mouse myeloma cell lines are employed. The hybridoma cells may be cultured in a suitable culture medium that preferably contains one or more substances that inhibit the growth or survival of the unfused, immortalized cells. For example, if the parental cells lack the enzyme hypoxanthine guanine phosphoribosyl transferase (HGPRT or HPRT), the culture medium for the hybridomas typically will include hypoxanthine, aminopterin, and thymidine ("HAT medium"), which substances prevent the growth of HGPRT-deficient cells.

Preferred immortalized cell lines are those that fuse efficiently, support stable high level expression of antibody by the selected antibody-producing cells, and are sensitive to a medium such as HAT medium. More preferred immortalized cell lines are murine myeloma lines, which can be obtained, for instance, from the Salk Institute Cell Distribution Center, San Diego, California and the American Type Culture Collection, Rockville, Maryland. Human myeloma and mouse-human heteromyeloma cell lines also have been described for the production of human monoclonal antibodies [Kozbor, J. Immunol., 133:3001 (1984); Brodeur et al., Monoclonal Antibody Production Techniques and Applications, Marcel Dekker, Inc., New York, (1987) pp. 51-63].

The culture medium in which the hybridoma cells are cultured can then be assayed for the presence of monoclonal antibodies directed against IL-11p. Preferably, the binding specificity of monoclonal antibodies produced by the hybridoma cells is determined by immunoprecipitation or by an *in vitro* binding assay, such as radioimmunoassay (RIA) or enzyme-linked immunosorbant assay (ELISA). Such techniques and assays are known in the art. The binding affinity of the monoclonal antibody can, for example, be determined by the Scatchard analysis of Munson and Pollard. *Anal. Biochem.*, 107:220 (1980).

After the desired hybridoma cells are identified, the clones may be subcloned by limiting dilution procedures and grown by standard methods [Goding, <u>supra</u>]. Suitable culture media for this purpose include, for example, Dulbecco's Modified Eagle's Medium and RPMI-1640 medium. Alternatively, the hybridoma cells may be grown in vivo as ascites in a mammal.

The monoclonal antibodies secreted by the subclones may be isolated or purified from the culture medium or ascites fluid by conventional immunoglobulin purification procedures such as, for example, protein A-Sepharose, hydroxylapatite chromatography, gel electrophoresis, dialysis, or affinity chromatography.

The monoclonal antibodies may also be made by recombinant DNA methods, such as those described in U.S. Patent No. 4,816,567. DNA encoding the monoclonal antibodies of the invention can be readily isolated and sequenced using conventional procedures (e.g., by using oligonucleotide probes that are capable of binding specifically to genes encoding the heavy and light chains of murine antibodies). The hybridoma cells of the invention serve as a preferred

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source of such DNA. Once isolated, the DNA may be placed into expression vectors, which are then transfected into host cells such as simian COS cells, Chinese hamster ovary (CHO) cells, or myeloma cells that do not otherwise produce immunoglobulin protein, to obtain the synthesis of monoclonal antibodies in the recombinant host cells. The DNA also may be modified, for example, by substituting the coding sequence for human heavy and light chain constant domains in place of the homologous murine sequences [U.S. Patent No. 4,816,567; Morrison et al., <a href="supra">supra</a>] or by covalently joining to the immunoglobulin coding sequence all or part of the coding sequence for a non-immunoglobulin polypeptide. Such a non-immunoglobulin polypeptide can be substituted for the constant domains of an antibody of the invention, or can be substituted for the variable domains of one antigen-combining site of an antibody of the invention to create a chimeric bivalent antibody.

The antibodies may be monovalent antibodies. Methods for preparing monovalent antibodies are well known in the art. For example, one method involves recombinant expression of immunoglobulin light chain and modified heavy chain. The heavy chain is truncated generally at any point in the Fc region so as to prevent heavy chain crosslinking. Alternatively, the relevant cysteine residues are substituted with another amino acid residue or are deleted so as to prevent crosslinking.

In vitro methods are also suitable for preparing monovalent antibodies. Digestion of antibodies to produce fragments thereof, particularly, Fab fragments, can be accomplished using routine techniques known in the art.

## Human and Humanized Antibodies

The anti-IL-1lp antibodies of the invention may further comprise humanized antibodies or human antibodies. Humanized forms of non-human (e.g., murine) antibodies are chimeric immunoglobulins, immunoglobulin chains or fragments thereof (such as Fv, Fab, Fab', F(ab')2 or other antigen-binding subsequences of antibodies) which contain minimal sequence derived from non-human immunoglobulin. Humanized antibodies include human immunoglobulins (recipient antibody) in which residues from a complementary determining region (CDR) of the recipient are replaced by residues from a CDR of a non-human species (donor antibody) such as mouse, rat or rabbit having the desired specificity, affinity and capacity. In some instances, Fv framework residues of the human immunoglobulin are replaced by corresponding non-human residues. Humanized antibodies may also comprise residues which are found neither in the recipient antibody nor in the imported CDR or framework sequences. In general, the humanized antibody will comprise substantially all of at least one, and typically two, variable domains, in which all or substantially all of the CDR regions correspond to those of a non-human immunoglobulin and all or substantially all of the FR regions are those of a human immunoglobulin consensus sequence. The humanized antibody optimally also will comprise at least a portion of an immunoglobulin constant region (Fc), typically that of a human immunoglobulin [Jones et al., Nature, 321: 522-525 (1986); Riechmann et al., Nature, 332: 323-329 (1988); and Presta, Curr. Op. Struct. Biol., 2: 593-596 (1992)].

Methods for humanizing non-human antibodies are well known in the art. Generally, a humanized antibody has one or more amino acid residues introduced into it from a source which

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is non-human. These non-human amino acid residues are often referred to as "import" residues, which are typically taken from an "import" variable domain. Humanization can be essentially performed following the method of Winter and co-workers [Jones et al., Nature, 321: 522-525 (1986); Riechmann et al., Nature, 332: 323-327 (1988); Verhoeyen et al., Science, 239: 1534-1536 (1988)], by substituting rodent CDRs or CDR sequences for the corresponding sequences of a human antibody. Accordingly, such "humanized" antibodies are chimeric antibodies (U.S. Patent No. 4,816,567), wherein substantially less than an intact human variable domain has been substituted by the corresponding sequence from a non-human species. In practice, humanized antibodies are typically human antibodies in which some CDR residues and possibly some FR residues are substituted by residues from analogous sites in rodent antibodies.

Human antibodies can also be produced using various techniques known in the art, including phage display libraries [Hoogenboom and Winter, J. Mol. Biol., 227: 381 (1991); Marks et al., J. Mol. Biol., 222: 581 (1991)]. The techniques of Cole et al. and Boerner et al. are also available for the preparation of human monoclonal antibodies (Cole et al., Monoclonal Antibodies and Cancer Therapy, Alan R. Liss, p. 77 (1985) and Boerner et al., J. Immunol., 147(1): 86-95 (1991)]. Similarly, human antibodies can be made by introducing of human immunoglobulin loci into transgenic animals, e.g., mice in which the endogenous immunoglobulin genes have been partially or completely inactivated. Upon challenge, human antibody production is observed, which closely resembles that seen in humans in all respects, including gene rearrangement, assembly, and antibody repertoire. This approach is described, for example, in U.S. Patent Nos. 5,545,807; 5,545,806; 5,569,825; 5,625,126; 5,633,425; 5,661,016, and in the following scientific publications: Marks et al., Bio/Technology 10, 779-783 (1992); Lonberg et al., Nature 368: 856-859 (1994); Morrison, Nature 368, 812-13 (1994); Fishwild et al., Nature Biotechnology 14, 845-51 (1996); Neuberger, Nature Biotechnology 14, 826 (1996); Lonberg and Huszar, Intern. Rev. Immunol. 13 65-93 (1995).

#### Bispecific Antibodies

Bispecific antibodies are monoclonal, preferably human or humanized, antibodies that have binding specificities for at least two different antigens. In the present case, one of the binding specificities is for the IL-1lp, the other one is for any other antigen, and preferably for a cell-surface protein or receptor or receptor subunit.

Methods for making bispecific antibodies are known in the art. Traditionally, the recombinant production of bispecific antibodies is based on the co-expression of two immunoglobulin heavy-chain/light-chain pairs, where the two heavy chains have different specificities [Milstein and Cuello, Nature, 305:537-539 (1983)]. Because of the random assortment of immunoglobulin heavy and light chains, these hybridomas (quadromas) produce a potential mixture of ten different antibody molecules, of which only one has the correct bispecific structure. The purification of the correct molecule is usually accomplished by affinity chromatography steps. Similar procedures are disclosed in WO 93/08829, published 13 May 1993, and in Traunecker et al., EMBO J., 10: 3655-3659 (1991).

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Antibody variable domains with the desired binding specificities (antibody-antigen combining sites) can be fused to immunoglobulin constant domain sequences. The fusion preferably is with an immunoglobulin heavy-chain constant domain, comprising at least part of the hinge, CH2, and CH3 regions. It is preferred to have the first heavy-chain constant region (CH1) containing the site necessary for light-chain binding present in at least one of the fusions. DNAs encoding the immunoglobulin heavy-chain fusions and, if desired, the immunoglobulin light chain, are inserted into separate expression vectors, and are co-transfected into a suitable host organism. For further details of generating bispecific antibodies see, for example, Suresh et al., Methods in Enzymology, 121:210 (1986).

### Heteroconjugate Antibodies

Heteroconjugate antibodies are also within the scope of the present invention. Heteroconjugate antibodies are composed of two covalently joined antibodies. Such antibodies have, for example, been proposed to target immune system cells to unwanted cells [U.S. Patent No. 4,676,980], and for treatment of HIV infection [WO 91/00360; WO 92/200373; EP 03089]. It is contemplated that the antibodies may be prepared *in vitro* using known methods in synthetic protein chemistry, including those involving crosslinking agents. For example, immunotoxins may be constructed using a disulfide exchange reaction or by forming a thioether bond. Examples of suitable reagents for this purpose include iminothiolate and methyl-4-mercaptobutyrimidate and those disclosed, for example, in U.S. Patent No. 4,676,980.

## E. Uses for anti-IL-11p Antibodies

The anti-IL-11p antibodies of the invention have various utilities. For example, anti-IL11p antibodies may be used in diagnostic assays for IL-11p, e.g., detecting its expression in
specific cells, tissues, or serum. Various diagnostic assay techniques known in the art may be
used, such as competitive binding assays, direct or indirect sandwich assays and
immunoprecipitation assays conducted in either heterogeneous or homogeneous phases [Zola,
Monoclonal Antibodies: A Manual of Techniques, CRC Press, Inc. (1987) pp. 147-158]. The
antibodies used in the diagnostic assays can be labeled with a detectable moiety. The detectable
moiety should be capable of producing, either directly or indirectly, a detectable signal. For
example, the detectable moiety may be a radioisotope, such as 3H, 14C, 32P, 35S, or 125I, a
fluorescent or chemiluminescent compound, such as fluorescein isothiocyanate, rhodamine, or
luciferin, or an enzyme, such as alkaline phosphatase, beta-galactosidase or horseradish
peroxidase. Any method known in the art for conjugating the antibody to the detectable moiety
may be employed, including those methods described by Hunter et al., Nature, 144: 945 (1962);
David et al., Biochemistru, 13: 1014 (1974); Pain et al., J. immunol. Meth., 40: 219 (1981); and
Nygren, J. Histochem. and Cytochem., 30: 407 (1982).

Anti-IL-1lp antibodies also are useful for the affinity purification of IL-1lp from recombinant cell culture or natural sources. In this process, the antibodies against IL-1lp are immobilized on a suitable support, such a Sephadex resin or filter paper, using methods known in the art. The immobilized antibody then is contacted with a sample containing the IL-1lp to be purified, and thereafter the support is washed with a suitable solvent that will remove

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substantially all the material in the sample except the IL-1lp, which is bound to the immobilized antibody. Finally, the support is washed with another suitable solvent that will release the IL-1lp from the antibody.

In addition, anti-IL-11p antibodies are useful as therapeutic agents for targeting of native IL-11p in IL-11p-mediated disease conditions, e.g. disease states characterized by pathologic IL-1 or IL-18 agonist or agonist-like activity of the native IL-11p. In the treatment and prevention of a native IL-11p-mediated disorder with the anti-IL-11p antibody of the invention, the antibody composition will be formulated, dosed, and administered in a fashion consistent with good medical practice. Factors for consideration in this context include the particular disorder being treated, the particular mammal being treated, the clinical condition of the individual patient, the cause of the disorder, the site of delivery of the antibody, the particular type of antibody, the method of administration, the scheduling of administration, and other factors known to medical practitioners. The "effective amount" or "therapeutically effective amount" of antibody to be administered will be governed by such considerations, and is the minimum amount necessary to prevent, ameliorate, or treat the native IL-11p-mediated disorder, including treating inflammatory diseases and reducing inflammatory responses. Such amount is preferably below the amount that is toxic to the host or renders the host significantly more susceptible to infections.

As a general proposition, the initial pharmaceutically effective amount of the antibody or antibody fragment administered parenterally per dose will be in the range of about 0.1 to 50~mg/kg of patient body weight per day, with the typical initial range of antibody used being 0.3~to~20~mg/kg/day, more preferably 0.3~to~15~mg/kg/day.

In one embodiment, using systemic administration, the initial pharmaceutically effective amount will be in the range of about 2 to 5 mg/kg/day.

For methods of the invention using administration by inhalation, the initial pharmaceutically effective amount will be in the range of about 1 microgram (ug)/kg/day to 100 mg/kg/day.

In one embodiment, the invention provides a method for treating an IL-11p-mediated inflammatory disorder comprising administering to a mammal, such as human, in need of such treatment an effective amount of an anti-IL-11p antibody.

In another embodiment, the invention provides a method for treating an hIL-Ilpmediated asthmatic disorder comprising administering to a human in need of such treatment an effective amount of an anti-hIL-Ilp antibody.

In another embodiment, the invention provides a method for treating an hIL-1Ra1-mediated asthmatic disorder comprising administering to a human in need of such treatment an effective amount of an anti-hIL-1Ra1 antibody.

In another embodiment, the invention provides a method for treating an IL-1lp-mediated rheumatoid arthritic disorder comprising administering to a mammal, such as human, in need of such treatment an effective amount of an anti-IL-1lp antibody.

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In another embodiment, the invention provides a method for treating an hIL-11pmediated rheumatoid arthritic disorder comprising administering to a human in need of such treatment an effective amount of an anti-hIL-11p antibody.

In another embodiment, the invention provides a method for treating an hlL-1Ra1mediated rheumatoid arthritic disorder comprising administering to a human in need of such treatment an effective amount of an anti-hlL-1Ra1 antibody.

In another embodiment, the invention provides a method for treating an IL-1lp-mediated osteoarthritic disorder comprising administering to a mammal, such as human, in need of such treatment an effective amount of an anti-IL-1lp antibody.

In another embodiment, the invention provides a method for treating an hIL-1lpmediated osteoarthritic disorder comprising administering to a human in need of such treatment an effective amount of an anti-hIL-1lp antibody.

In another embodiment, the invention provides a method for treating an hIL-1Ra1mediated osteoarthritic disorder comprising administering to a human in need of such treatment an effective amount of an anti-hIL-1Ra1 antibody.

In another embodiment, the invention provides a method for treating an IL-11p-mediated septic disorder comprising administering to a mammal, such as human, in need of such treatment an effective amount of an anti-IL-11p antibody.

In another embodiment, the invention provides a method for treating an hIL-1lpmediated septic disorder comprising administering to a human in need of such treatment an effective amount of an anti-hIL-1lp antibody.

In another embodiment, the invention provides a method for treating an hIL-1Ra1mediated septic disorder comprising administering to a human in need of such treatment an effective amount of an anti-hIL-1Ra1 antibody.

In another embodiment, the invention provides a method for treating IL-1lp-mediated acute lung injury comprising administering to a mammal, such as human, in need of such treatment an effective amount of an anti-IL-1lp antibody.

In another embodiment, the invention provides a method for treating hIL-llp-mediated acute lung injury comprising administering to a human in need of such treatment an effective amount of an anti-hIL-llp antibody.

In another embodiment, the invention provides a method for treating hIL-1Ra1-mediated acute lung injury comprising administering to a human in need of such treatment an effective amount of an anti-hIL-1Ra1 antibody.

In another embodiment, the invention provides a method for treating IL-1lp-mediated adult respiratory distress syndrome comprising administering to a mammal, such as human, in need of such treatment an effective amount of an anti-IL-1lp antibody.

In another embodiment, the invention provides a method for treating hIL-11p-mediated adult respiratory distress syndrome comprising administering to a human in need of such treatment an effective amount of an anti-hIL-11p antibody.

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In another embodiment, the invention provides a method for treating hIL-1Ra1-mediated adult respiratory distress syndrome comprising administering to a human in need of such treatment an effective amount of an anti-hIL-1Ra1 antibody.

In another embodiment, the invention provides a method for treating IL-1lp-mediated idiopathic pulmonary fibrosis comprising administering to a mammal, such as human, in need of such treatment an effective amount of an anti-IL-1lp antibody.

In another embodiment, the invention provides a method for treating hIL-1lp-mediated idiopathic pulmonary fibrosis comprising administering to a human in need of such treatment an effective amount of an anti-hlL-1lp antibody.

In another embodiment, the invention provides a method for treating hIL-1Ra1-mediated idiopathic pulmonary fibrosis comprising administering to a human in need of such treatment an effective amount of an anti-hIL-1Ra1 antibody.

In another embodiment, the invention provides a method for treating an IL-11p-mediated ischemic reperfusion disease, such as surgical tissue reperfusion injury, stroke, myocardial ischemia, or acute myocardial infarction, comprising administering to a mammal, such as human, in need of such treatment an effective amount of an anti-IL-1lp antibody.

In another embodiment, the invention provides a method for treating an hIL-11pmediated ischemic reperfusion disease, such as surgical tissue reperfusion injury, stroke, myocardial ischemia, or acute myocardial infarction, comprising administering to a human in need of such treatment an effective amount of an anti-hIL-11p antibody.

In another embodiment, the invention provides a method for treating an hIL-1Ra1mediated ischemic reperfusion disease, such as surgical tissue reperfusion injury, stroke, myocardial ischemia, or acute myocardial infarction, comprising administering to a human in need of such treatment an effective amount of an anti-hIL-1Ra1 antibody.

In another embodiment, the invention provides a method for treating an IL-11p-mediated psoriatic disorder comprising administering to a mammal, such as human, in need of such treatment an effective amount of an anti-IL-11p antibody.

In another embodiment, the invention provides a method for treating an hIL-11pmediated psoriatic disorder comprising administering to a human in need of such treatment an effective amount of an anti-hIL-11p antibody.

In another embodiment, the invention provides a method for treating an hIL-1Ra1mediated psoriatic disorder comprising administering to a human in need of such treatment an effective amount of an anti-hIL-1Ra1 antibody.

In another embodiment, the invention provides a method for treating an IL-11p-mediated graft-versus-host disease (GVHD) comprising administering to a mammal, such as human, in need of such treatment an effective amount of an anti-IL-11p antibody.

In another embodiment, the invention provides a method for treating an hIL-llpmediated graft-versus-host disease (GVHD) comprising administering to a human in need of such treatment an effective amount of an anti-hIL-1lp antibody.

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In another embodiment, the invention provides a method for treating an hIL-1Ra1-mediated graft-versus-host disease (GVHD) comprising administering to a human in need of such treatment an effective amount of an anti-hIL-1Ra1 antibody.

In another embodiment, the invention provides a method for treating an IL-11p-mediated inflammatory bowel disease such as ulcerative colitis, comprising administering to a mammal, such as human, in need of such treatment an effective amount of an anti-IL-11p antibody.

In another embodiment, the invention provides a method for treating an hIL-11p-mediated inflammatory bowel disease such as ulcerative colitis, comprising administering to a human in need of such treatment an effective amount of an anti-hIL-11p antibody.

In another embodiment, the invention provides a method for treating an hIL-1Ra1-mediated inflammatory bowel disease such as ulcerative colitis, comprising administering to a human in need of such treatment an effective amount of an anti-hIL-1Ra1 antibody.

The following examples are offered for illustrative purposes only, and are not intended to limit the scope of the present invention in any way.

All patent and literature references cited in the present specification are hereby incorporated by reference in their entirety.

## EXAMPLES

Commercially available reagents referred to in the examples were used according to manufacturer's instructions unless otherwise indicated. The source of those cells identified in the following examples, and throughout the specification, by ATCC accession numbers is the American Type Culture Collection, Manassas, VA.

#### EXAMPLE 1

# Isolation of DNA encoding hIL-1Ra1 and mlL-1Ra3

A public expressed sequence tag (EST) DNA database (Genbank) was searched with human interleukin-1 receptor antagonist (hIL-1Ra) sequence, also known as secretory human interleukin-1 receptor antagonist ('sIL-1Ra') sequence, and a human EST designated Al014548 (which corresponds to nucleotides 145-629 of Figure 4, SEQ ID NO:8), and a murine EST designated W08205 (Figure 10, SEQ ID NO:17), were identified, which showed homology with the known protein hIL-1Ra (sIL-1Ra).

EST clones Al014548 and W08205 were purchased from Research Genetics (Huntsville, AL) and the cDNA inserts were obtained and sequenced in their entireties.

The entire nucleotide sequence of the clone Al014548, designated DNA85066, is shown in Figure 1 (SEQ ID NO:1). Clone DNA85066 contains a single open reading frame that is interrupted by an apparent intronic sequence. The intron is bounded by splice junctions at nucleotide positions 181 to 186 (splice donor site) and nucleotide positions 430 to 432 (splice acceptor site) (Fig.1; SEQ ID NO:1).

A virtual processed nucleotide sequence (Fig. 3; SEQ ID NO:6), designated DNA94618, was derived by removing the apparent intronic sequence from clone DNA85066. Clone DNA94618 contains a single open reading frame with an apparent translational initiation site at nucleotide positions 103-105, and a stop codon at nucleotide positions 682-684 (Fig. 3; SEQ ID NO:6). The predicted polypeptide precursor (hIL-IRa1) (Fig. 3; SEQ ID NO:7) is 193 amino acids long. The putative signal sequence extends from amino acid positions 1 to 14. A putative

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cAMP- and cGMP-dependent protein kinase phosphorylation site is located at amino acid positions 33-36. Putative N-myristoylation sites are located at amino acid positions 50-55 and 87-92.

Clone DNA85066 (designated as DNA85066-2534) has been deposited with ATCC and was assigned ATCC deposit no. 203588. The full-length hIL-1Ra1 protein shown in Figure 3 (SEQ ID NO:7) has an estimated molecular weight of about 21,822 daltons and a pl of about 8.9.

Based on a sequence alignment analysis of the full-length sequence (SEQ ID NO:7), hIL-1Ra1 shows significant amino acid sequence identity to hIL-1Ra (sIL-1Ra) and hIL-1Ra $\beta$  proteins.

The entire nucleotide sequence of the clone W08205, designated DNA92505, is shown in Figure 9 (SEQ ID NO:15). Clone DNA92505 contains a single open reading frame with an apparent translational initiation site at nucleotide positions 145-147, and a stop codon at nucleotide positions 610-612 (Fig. 9; SEQ ID NO:15). The predicted polypetide precursor (mIL-1Ra3) (Fig. 9; SEQ ID NO:16) is 155 amino acids long. The putative signal sequence extends from amino acid positions 1-33. Putative N-myristoylation sites are located at amino acid positions 29-34, 60-65, 63-68, 91-96 and 106-111. An interleukin-1-like sequence is located at amino acid positions 111-131.

Clone DNA92505 (designated as DNA92505-2534) was deposited with ATCC and was assigned ATCC deposit no. 203590. The full length mIL-1 Ra3 protein shown in Figure 9 (SEQ ID NO:16) has an estimated molecular weight of about 17,134 daltons and a pl of about 4.8.

Based on a sequence alignment analysis of the full-length sequence (SEQ ID NO:16), mlL-1Ra3 shows significant amino acid sequence identity to mIL-1Ra, hiclL-1Ra, hIL-1Ra (sIL-1Ra) and hIL-1Ra8 proteins.

### EXAMPLE 2

## Isolation of DNA encoding hlL-1ra2 and hlL-1Ra3

A expressed sequence tag (EST) DNA database (LIFESEQ®, Incyte Pharmaceuticals, Palo Alto, CA) was searched with human interleukin-1 receptor antagonist (hlL-1Ra) sequence, also known as secretory human interleukin-1 receptor antagonist ("slL-1Ra") sequence, and the ESTs, designated 1433156 (Figure 5, SEQ ID NO:11) and 5120028 (Figure 8, SEQ ID NO:14), were identified, which showed homology with the hlL-1Ra known protein.

EST clones 1433156 and 5120028 were purchased from Incyte Pharmaceuticals (Palo Alto, CA) and the cDNA inserts were obtained and sequenced in their entireties.

The entire nucleotide sequence of the clone 1433156, designated DNA92929, is shown in Figure 5 (SEQ ID NO:9). Clone DNA92929 contains a single open reading frame with an apparaturanslational initiation site at nucleotide positions 96-98, and a stop codon at nucleotide positions 498-500 (Fig. 5; SEQ ID NO:9). The predicted polypeptide precursor (hIL-1Ra2) (Fig. 5; SEQ ID NO:10) is 134 amino acids long. A putative signal sequence extends from amino acid positions 1-26.

Clone DNA92929 (designated as DNA92929-2534) was deposited with ATCC and was assigned ATCC deposit no. 203586. The full-length hIL-1Ra2 protein shown in Figure 5 (SEQ ID NO:10) has an estimated molecular weight of about 14,927 daltons and a pl of about 4.8.

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Based on a sequence alignment analysis of the full-length sequence (SEQ ID NO:10), hill-1Ra2 shows significant amino acid sequence identity to hlL-1Ra6 protein. hlL-1Ra2 is believed to be a splice variant of hlL-1Ra6.

The entire nucleotide sequence of the clone 5120028, designated DNA96787, is shown in Figure 7 (SEQ ID NO:12). Clone DNA96787 contains a single open reading frame with an apparent translational initiation site at nucleotide positions 1-3, and a stop codon at nucleotide positions 466-468 (Fig. 7; SEQ ID NO:12). The predicted polypeptide precursor (hIL-1Ra3) (Fig. SEQ ID NO:13) is 155 amino acids long. A putative signal sequence extends from amino acid positions 1-33. Putative N-myristoylation sites are located at amino acid positions 29-34, 60-65, 63-68, 73-78, 91-96 and 106-111. An interleukin-1-like sequence is located at amino acid positions 111-131.

It is believed that the predicted 155 amino acid polypeptide of hIL-1Ra3 behaves as a mature sequence (without a presequence that is removed in post-translational processing in certain animal cells. It is also believed that other animal cells recognize and remove one or more signal peptide(s) extending from amino acid positions 1 to about 33. As shown in Example 14 below, transiently transfected CHO host cells secrete a form of hIL-1Ra3 that only lacks the N-terminal methionine in the sequence of Figure 7 (SEQ ID NO:13).

Clone DNA96787 (designated as DNA96787-2534) was deposited with ATCC and was assigned ATCC deposit no. 203589. The full length hIL-1Ra3 protein shown in Figure 7 (SEQ ID NO:13) has an estimated molecular weight of about 16,961 daltons and a pI of about 4.9.

Based on a sequence alignment analysis of the full-length sequence (SEQ ID NO:13), hIL-1Ra3 shows significant amino acid sequence identity to hiclL-1Ra and hlL-1Ra (sIL-1Ra) proteins.

## EXAMPLE 3

## Northern Blot Analysis

Expression of hIL-1Ra3 mRNA in human tissues and mIL-1Ra3 mRNA in mouse tissues was examined by Northern blot analysis. Human and mouse multiple tissue northern (RNA) blots and mouse embryo blots were purchased from Clontech and probed with corresponding cDNA according to the manufacturer's instructions.

As shown in Fig. 11, hIL-1Ra3 mRNA (2.7 kb) were detected only in human placenta and mIL-1Ra3 mRNA transcripts (1.4 kb and 2.5 kb) were detected only in the day-17 mouse embryo.

EXAMPLE 4

#### Use of IL-11p as a hybridization probe

The following method describes use of a nucleotide sequence encoding IL-1lp as a hybridization probe.

DNA comprising the coding sequence of full-length IL-1lp (as shown in Figures 3, 5, 7, 9, 15, 16 and 19; SEQ ID NOS:6, 9, 12, 15, 18, 20 and 24) is employed as a probe to screen for homologous DNAs (such as those encoding naturally-occurring variants of IL-1lp) in human tissue cDNA libraries or human tissue genomic libraries.

Hybridization and washing of filters containing either library DNAs is performed under the following high stringency conditions. Hybridization of radiolabeled IL-11p-derived probe to

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the filters is performed in a solution of 50% formamide, 5x SSC, 0.1% SDS, 0.1% sodium pyrophosphate, 50 mM sodium phosphate, pH 6.8, 2x Denhardt's solution, and 10% dextran sulfate at 42°C for 20 hours. Washing of the filters is performed in an aqueous solution of 0.1x SSC and 0.1% SDS at 42°C.

DNAs having a desired sequence identity with the DNA encoding full-length native sequence IL-11p can then be identified using standard techniques known in the art.

#### EXAMPLE 5

## Expression of IL-11p in E. coli

This example illustrates preparation of an unglycosylated form of IL-1lp by recombinant expression in E. coli.

The DNA sequence encoding an IL-1lp is initially amplified using selected PCR primers. The primers should contain restriction enzyme sites which correspond to the restriction enzyme sites on the selected expression vector. A variety of expression vectors may be employed. An example of a suitable vector is pBR322 (derived from E. coli; see Bolivar et al., Gene, 2:95 (1977)) which contains genes for ampicillin and tetracycline resistance. The vector is digested with restriction enzyme and dephosphorylated. The PCR amplified sequences are then ligated into the vector. The vector will preferably include sequences which encode for an antibiotic resistance gene, a trp promoter, a polyhis leader (including the first six STII codons, polyhis sequence, and enterokinase cleavage site), the IL-1lp coding region, lambda transcriptional terminator, and an argU gene.

The ligation mixture is then used to transform a selected *E. coli* strain using the methods described in Sambrook et al., <u>supra</u>. Transformants are identified by their ability to grow on LB plates and antibiotic resistant colonies are then selected. Plasmid DNA can be isolated and confirmed by restriction analysis and DNA sequencing.

Selected clones can be grown overnight in liquid culture medium such as LB broth supplemented with antibiotics. The overnight culture may subsequently be used to inoculate a larger scale culture. The cells are then grown to a desired optical density, during which the expression promoter is turned on.

After culturing the cells for several more hours, the cells can be harvested by centrifugation. The cell pellet obtained by the centrifugation can be solubilized using various agents known in the art, and the solubilized IL-1lp protein can then be purified using a metal chelating column under conditions that allow tight binding of the protein.

#### EXAMPLE 6

## Expression of IL-11p in mammalian cells

This example illustrates preparation of a potentially glycosylated form of IL-1lp by recombinant expression in mammalian cells.

The vector, pRK5 (see EP 307,247, published March 15, 1989), is employed as the expression vector. Optionally, the IL-1lp DNA is ligated into pRK5 with selected restriction enzymes to allow insertion of the IL-1lp DNA using ligation methods such as described in Sambrook et al., supra. The resulting vector is called pRK5-IL-1lp.

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In one embodiment, the selected host cells may be 293 cells. Human 293 cells (ATCC CCL 1573) are grown to confluence in tissue culture plates in medium such as DMEM supplemented with fetal calf serum and optionally, nutrient components and/or antibiotics. About 10 µg pRK5-IL-11p DNA is mixed with about 1 µg DNA encoding the VA RNA gene [Thimmappaya et al., Cell, 31:543 (1982)] and dissolved in 500 µl of 1 mM Tris-HCl, 0.1 mM EDTA, 0.227 M CaCl<sub>2</sub>. To this mixture is added, dropwise, 500 µl of 50 mM HEPES (pH 7.35), 280 mM NaCl, 1.5 mM NaPO<sub>4</sub>, and a precipitate is allowed to form for 10 minutes at 25°C. The precipitate is suspended and added to the 293 cells and allowed to settle for about four hours at 37°C. The culture medium is aspirated off and 2 ml of 20% glycerol in PBS is added for 30 seconds. The 293 cells are then washed with serum free medium, fresh medium is added and the cells are incubated for about 5 days.

Approximately 24 hours after the transfections, the culture medium is removed and replaced with culture medium (alone) or culture medium containing 200  $\mu$ Ci/ml <sup>38</sup>S-cysteine and 200  $\mu$ Ci/ml <sup>38</sup>S-methionine. After a 12 hour incubation, the conditioned medium is collected, concentrated on a spin filter, and loaded onto a 15% SDS gel. The processed gel may be dried and exposed to film for a selected period of time to reveal the presence of IL-1lp polypeptide. The cultures containing transfected cells may undergo further incubation (in serum free medium) and the medium is tested in selected bioassays.

In an alternative technique, IL-11p may be introduced into 293 cells transiently using the dextran sulfate method described by Somparyrac et al., Proc. Natl. Acad. Sci. USA, 12: 7575 (1981). 293 cells are grown to maximal density in a spinner flask and 700 µg pRK5-IL-11p DNA is added. The cells are first concentrated from the spinner flask by centrifugation and washed with PBS. The DNA-dextran precipitate is incubated on the cell pellet for four hours. The cells are treated with 20% glycerol for 90 seconds, washed with tissue culture medium, and reintroduced into the spinner flask containing tissue culture medium, 5 µg/ml bovine insulin and 0.1 µg/ml bovine transferrin. After about four days, the conditioned media is centrifuged and filtered to remove cells and debris. The sample containing expressed IL-11p can then be concentrated and purified by any selected method, such as dialysis and/or column chromatography.

In another embodiment, IL-1lp can be expressed in CHO cells. The pRK5-IL-1lp can be transfected into CHO cells using known reagents such as CaPO<sub>4</sub> or DEAE-dextran. As described above, the cell cultures can be incubated, and the medium replaced with culture medium (alone) or medium containing a radiolabel such as <sup>38</sup>S-methionine. After determining the presence of IL-1lp polypeptide, the culture medium may be replaced with serum free medium. Preferably, the cultures are incubated for about 6 days, and then the conditioned medium is harvested. The medium containing the expressed IL-1lp can then be concentrated and purified by any selected method.

Epitope-tagged IL-11p may also be expressed in host CHO cells. The IL-11p may be subcloned out of the pRK5 vector. The subclone insert can undergo PCR to fuse in frame with a selected epitope tag such as a poly-his tag into a Baculovirus expression vector. The poly-his

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tagged IL-1lp insert can then be subcloned into a SV40 driven vector containing a selection marker such as DHFR for selection of stable clones. Finally, the CHO cells can be transfected (as described above) with the SV40 driven vector. Labeling may be performed, as described above, to verify expression. The culture medium containing the expressed poly-His tagged IL-1lp can then be concentrated and purified by any selected method, such as by Ni<sup>2+</sup>-chelate affinity chromatography.

### EXAMPLE 7

## Expression of IL-1lp in Yeast

The following method describes recombinant expression of IL-11p in yeast.

First, yeast expression vectors are constructed for intracellular production or secretion of IL-1lp from the ADH2/GAPDH promoter. DNA encoding IL-1lp and the promoter is inserted into suitable restriction enzyme sites in the selected plasmid to direct intracellular expression of IL-1lp. For secretion, DNA encoding IL-1lp can be cloned into the selected plasmid, together with DNA encoding the ADH2/GAPDH promoter, a native IL-1lp signal peptide or other mammalian signal peptide, or, for example, a yeast alpha-factor or invertase secretory signal/leader sequence, and linker sequences (if needed) for expression of IL-1lp.

Yeast cells, such as yeast strain AB110, can then be transformed with the expression plasmids described above and cultured in selected fermentation media. The transformed yeast supernatants can be analyzed by precipitation with 10% trichloroacetic acid and separation by SDS-PAGE, followed by staining of the gels with Coomassie Blue stain.

Recombinant IL-1lp can subsequently be isolated and purified by removing the yeast cells from the fermentation medium by centrifugation and then concentrating the medium using selected cartridge filters. The concentrate containing IL-1lp may further be purified using selected column chromatography resins.

#### EXAMPLE 8

## Expression of IL-11p in Baculovirus-Infected Insect Cells

The following method describes recombinant expression of IL-11p in Baculovirus-infected insect cells.

The sequence coding for IL-1lp is fused upstream of an epitope tag contained within a baculovirus expression vector. Such epitope tags include poly-his tags and immunoglobulin tags (like Fc regions of IgG). A variety of plasmids may be employed, including plasmids derived from commercially available plasmids such as pVL1393 (Novagen). Briefly, the sequence encoding IL-1lp or the desired portion of the coding sequence of IL-1lp (such as the sequence encoding the mature protein) is amplified by PCR with primers complementary to the 5' and 3' regions. The 5' primer may incorporate flanking (selected) restriction enzyme sites. The product is then digested with those selected restriction enzymes and subcloned into the expression vector.

Recombinant baculovirus is generated by co-transfecting the above plasmid and BaculoGold™ virus DNA (Pharmingen) into Spodoptera frugiperda ("SI9") cells (ATCC CRL 1711) using lipofectin (commercially available from GIBCO-BRL). After 4 - 5 days of incubation at 28°C, the released viruses are harvested and used for further amplifications. Viral infection and

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protein expression are performed as described by O'Reilley et al., Baculovirus expression vectors:

A Laboratory Manual, Oxford: Oxford University Press (1994).

Expressed poly-his tagged IL-1lp can then be purified, for example, by Ni2+-chelate affinity chromatography as follows. Extracts are prepared from recombinant virus-infected Sf9 cells as described by Rupert et al., Nature, 362: 175-179 (1993). Briefly, Sf9 cells are washed, resuspended in sonication buffer (25 mL Hepes, pH 7.9; 12.5 mM MgCl<sub>2</sub>; 0.1 mM EDTA; 10% glycerol; 0.1% NP-40; 0.4 M KCl), and sonicated twice for 20 seconds on ice. The sonicates are cleared by centrifugation, and the supernatant is diluted 50-fold in loading buffer (50 mM phosphate, 300 mM NaCl, 10% glycerol, pH 7.8) and filtered through a 0.45  $\mu m$  filter. A Ni<sup>2</sup>\*-NTA agarose column (commercially available from Qiagen) is prepared with a bed volume of 5 mL, washed with 25 mL of water and equilibrated with 25 mL of loading buffer. The filtered cell extract is loaded onto the column at 0.5 mL per minute. The column is washed to baseline A280 with loading buffer, at which point fraction collection is started. Next, the column is washed with a secondary wash buffer (50 mM phosphate; 300 mM NaCl, 10% glycerol, pH 6.0), which elutes nonspecifically bound protein. After reaching A280 baseline again, the column is developed with a 0 to 500 mM Imidazole gradient in the secondary wash buffer. One mL fractions are collected and analyzed by SDS-PAGE and silver staining or Western blot with Ni2+-NTAconjugated to alkaline phosphatase (Qiagen). Fractions containing the eluted His10-tagged IL-11p are pooled and dialyzed against loading buffer.

Alternatively, purification of the IgG tagged (or Fc tagged) IL-1lp can be performed using known chromatography techniques, including for instance, Protein A or protein G column chromatography.

## EXAMPLE 9

# IL-18 Receptor and IL-1 Receptor Binding of hIL-1Ra1

To facilitate the characterization of hIL-1Ra1, a PCR fragment containing the partial ORF of cione DNA85066 (Figure 1; SEQ ID NO:3) was cloned into pCMV1FLAG (IBI Kodak, described in Pan et al., Science, 276: 111-113) as an in-frame fusion to a NH<sub>2</sub>-terminal preprotrypsin leader sequence and FLAG tag encoded by the vector. The entire cDNA insert of the recombinant pCMV1FLAG vector clone (designated clone DNA96786) was sequenced (Figure 2; SEQ ID NO:4). The cDNAs encoding the extracellular domain of hIL1R and hIL18R (formerly known as hIL1Rrp) were obtained by polymerase chain reaction (PCR) and cloned into a modified pCMV1FLAG vector that allowed for in-frame fusion with the Fc portion of human immunoglobulin G.

Human embryonic kidney 293 cells were grown in high glucose DMEM (Genentech, Inc). The cells were seeded at 3-4 X10° per plate (100 mm) and co-transfected with pCMV1FLAG-IIL-1Ra1 and pCMV1FLAG-ILIR-ECD-Fc or pCMV1FLAG-IL18R-ECD-Fc by means of calcium phosphate precipitation. The media were changed 12 hours post transfection. The resultant conditioned media (10 ml each) were harvested after a further 70-74 hour incubation, clarified by centrifugation, aliquoted and stored at -70°C. The receptor-Fc and ligand complex from 1.5 ml conditioned medium was immunoprecipitated with protein G-Sepharose, washed three times with buffer containing 50 mM Hepes, pH7.0, 150 mM NaCl, 1 mM EDTA, 1% NP-40, and a

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protease inhibitor cocktail (BMB) and resolved on a 10-20% SDS-PAGE gel. The bound ligand was identified by immunoblotting using anti-FLAG monoclonal antibody (BMB).

As shown in Figure 13A, the secreted FLAGhIL-1Ra1 fusion protein bound to IL-18R ECD and did not bind to IL-1R ECD, which indicates that hIL-1Ra1 could be an agonist or antagonist of IL-18R.

#### EXAMPLE 10

## IL-1 Receptor and IL-18 Receptor Binding of mIL-1Ra3 and hlL-1Ra3

cDNA encoding mIL-1Ra3 (DNA92505 shown in Figure 9; SEQ ID NO:15) was cloned into pRK7 with a carboxy-terminal FLAG-tag. The resulting expression construct was transfected into human embryonic kidney 293 cells by means of calcium phosphate precipitation. 84-90 hours post transfection, the conditioned media containing secreted FLAGmIL-1Ra3 fusion protein was harvested. Conditioned media containing secreted IL-18R-Fc and IL-1R-Fc proteins were prepared as described in Example 9 above, with the exception that the 293 cells were transfected with either pCMV1FLAG-IL1R-ECD-Fc or pCMV1FLAG-IL18R-ECD-Fc alone (without pCMV1FLAG-IL-1Ra1 cotransfection).

For in vitro binding assays, IL-1R-Fc or IL-18R-Fc from 0.5 ml of the conditioned medium was immobilized to protein G-agarose and then mixed with 1.2 ml conditioned medium containing FLAGmIL-1Ra3. The receptor-ligand complexes were washed and resolved on an 10-20% SDS-PAGE gel and the bound ligand was detected by immunobloting using anti-FLAG monoclonal antibody (Boehringer Mannheim).

As shown in Figure 14, FLAGmIL-1Ra3 fusion protein bound to IL-1R ECD and did not bind to IL-18R ECD. Since the amino acid sequence of mIL-1Ra3 is related to that of the known interleukin-1 receptor antagonist protein (IL-1Ra), mIL-3Ra3 is believed to be a novel IL-1 receptor antagonist.

cDNA encoding hIL-1Ra3 (DNA96787 shown in Figure 7; SEQ ID NO:12) was cloned into pRK7 with a carboxy-terminal FLAG tag to form pRK7hIL-1Ra3-FLAG. pCMV1FLAG-IL1R-ECD-Fc and pCMV1FLAG-IL18R-ECD-Fc were obtained as described in Example 9 above. Similarly, cDNA encoding DR6 was cloned into the modified pCMV1FLAG vector of Example 9 to form pCMV1FLAG-DR6-Fc, encoding DR6 fused to the Fc portion of human immunoglobulin G. Conditioned media containing (1) a combination of secreted FLAGhIL-1Ra3 and FLAG-DR6-Fc (2) a combination of secreted FLAGhIL-1Ra3 and FLAG-IL1R-ECD-Fc or (3) a combination of secreted FLAGhIL-1Ra3 and FLAG-IL18R-ECD-Fc were prepared by cotransfecting Human 293 cells with (1) pRK7hIL-1Ra3-FLAG and pCMV1FLAG-DR6-Fc (2) pRK7hIL-1Ra3-FLAG and pCMV1FLAG-IL1R-ECD-Fc or (3) pRK7hIL-1Ra3-FLAG and pCMV1FLAG-IL18R-ECD-Fc, culturing the transfectant cells and harvesting culture media essentially as described in Example The receptor-Fc and ligand complex from each conditioned medium was 9 above. immunoprecipitated with protein G-Sepharose or anti-FLAG monoclonal antibody, and immunoprecipitates were resolved by gel electrophoresis and immunoblotting with anti-FLAG monoclonal antibody essentially as described in Example 9 above.

As shown in Figure 13B, FLAGhIL-1Ra3 fusion protein bound to IL-1R-ECD-Fc and did not bind to IL-18R-ECD-Fc or DR6-Fc. Since the amino acid sequence of hIL-1Ra3 is related to

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that of the known interleukin-1 receptor antagonist protein (IL-1Ra), hlL-3Ra3 is believed to be a novel IL-1 receptor antagonist.

## EXAMPLE 11

## Preparation of Antibodies that Bind IL-11p

This example illustrates preparation of monoclonal antibodies which can specifically bind IL-11p.

Techniques for producing the monoclonal antibodies are known in the art and are described, for instance, in Goding, supra. Immunogens that may be employed include purified IL-11p, fusion proteins containing IL-11p, and cells expressing recombinant IL-11p on the cell surface. Selection of the immunogen can be made by the skilled artisan without undue experimentation.

Mice, such as Balb/c, are immunized with the IL-1lp immunogen emulsified in complete Freund's adjuvant and injected subcutaneously or intraperitoneally in an amount from 1-100 micrograms. Alternatively, the immunogen is emulsified in MPL-TDM adjuvant (Ribi Immunochemical Research, Hamilton, MT) and injected into the animal's hind foot pads. The immunized mice are then boosted 10 to 12 days later with additional immunogen emulsified in the selected adjuvant. Thereafter, for several weeks, the mice may also be boosted with additional immunization injections. Serum samples may be periodically obtained from the mice by retro-orbital bleeding for testing in ELISA assays to detect anti-IL-1lp antibodies.

After a suitable antibody titer has been detected, the animals "positive" for antibodies can be injected with a final intravenous injection of IL-11p. Three to four days later, the mice are sacrificed and the spleen cells are harvested. The spleen cells are then fused (using 35% polyethylene glycol) to a selected murine myeloma cell line such as P3X63AgU.1, available from ATCC, No. CRL 1597. The fusions generate hybridoma cells which can then be plated in 96 well tissue culture plates containing HAT (hypoxanthine, aminopterin, and thymidine) medium to inhibit proliferation of non-fused cells, myeloma hybrids, and spleen cell hybrids.

The hybridoma cells will be screened in an ELISA for reactivity against IL-1lp. Determination of "positive" hybridoma cells secreting the desired monoclonal antibodies against IL-1lp is within the skill in the art.

The positive hybridoma cells can be injected intraperitoneally into syngeneic Balb/c mice to produce ascites containing the anti-IL-11p monoclonal antibodies. Alternatively, the hybridoma cells can be grown in tissue culture flasks or roller bottles. Purification of the monoclonal antibodies produced in the ascites can be accomplished using ammonium sulfate precipitation, followed by gel exclusion chromatography. Alternatively, affinity chromatography based upon binding of antibody to protein A or protein G can be employed.

### EXAMPLE 12

## Isolation of DNA encoding hIL-1Ra1L, hIL-1Ra1V and hIL-1Ra1S

Several intron-containing cDNA clones related to the hIL-1Ra1 intron-containing clone DNA85066 (Figure 2) (SEQ ID NO:4) were isolated from a human testis cDNA library and fully sequenced. The intron-containing cDNA sequences were used to determine a full-length open reading frame (ORF) with the GENESCAN program (http://CCR-

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081.mit.edu/GENESCAN.html). The ORF-encoding sequence was used to design two DNA primers, ggc gga tcc aaa atg ggc tct gag gac tgg g (SEQ ID NO:29) (1Ra1016) and gcg gaa ttc taa tcg ctg acc tca ctg ggg (SEO ID NO:30) (1Ra1017). The 1Ra1016 and 1Ra1017 primers were synthesized and used to clone cDNA from human fetal skin and SK-lu-1 cell cDNA libraries using polymerase chain reaction (PCR). Several PCR products were isolated and sequenced. Two full length cDNA clones (designated DNA102043 and DNA102044) from PCR products were found to encode hIL-1Ra1 isoforms.

The entire nucleotide sequence of clone DNA102043 is shown in Figure 15 (SEQ ID NO:18). Clone DNA102043 contains a single open reading frame with an apparent translational initiation site at nucleotide positions 4-6, and a stop codon at nucleotide positions 625-627 (Figure 15; SEQ ID NO:18). The predicted polypeptide precursor (designated hIL-1Ra1L) (Fig. 15; SEQ ID NO:19) is 207 amino acids long. The putative signal sequence extends from amino acid positions 1-34.

Clone DNA102043 (designated DNA102043-2534) was deposited with ATCC and was assigned ATCC deposit no. 203846. The full-length hlL-1Ra1L protein shown in Figure 15 (SEQ ID NO:19) has an estimated molecular weight of about 23,000 daltons and a pl of about 6.08.

Based on a sequence alignment analysis of the full length sequence (SEQ ID NO:19), hIL-1Ra1L shows significant amino acid sequence identity to hIL-1Raß and TANGO-77 protein. In addition, a portion of the DNA sequence of clone DNA102043 (Figure 15) (SEQ ID NO:18) was found to coincide with the DNA sequence of EST AI014548 (Figure 4) (SEQ ID NO:8) and with the complement of the DNA sequence of EST AI343258 (Figure 17) (SEQ ID NO:23).

The entire nucleotide sequence of clone DNA102044 is shown in Figure 16 (SEQ ID NO:20). Clone DNA102044 contains a single open reading frame with an apparent translational initiation site at nucleotide positions 4-6, and a stop codon at nucleotide positions 505-507 (Figure 16; SEQ ID NO:20). The predicted polypeptide (designated hIL-1Ra1S) (Fig. 16; SEQ ID NO:21) is 167 amino acids long, and it is believed to behave as a mature sequence (without a presequence that is removed in post-translational processing) in certain animal cells. In addition, it is believed that other animal cells recognize and remove in post-translational processing one or more signal peptide(s) contained in the sequence extending from amino acid positions 1 to about 46.

Clone DNA102044 (designated DNA102044-2534) was deposited with ATCC and was assigned ATCC deposit no. 203855. The full-length hIL-1Ra1S protein shown in Figure 16 (SEQ ID NO:21) has an estimated molecular weight of about 18,478 daltons and a pl of about 5.5.

Based on a sequence alignment analysis of the full length sequence (SEQ ID NO:21), hlL-1Ra1S appears to be an allelic variant of TANGO-77 protein and also shows significant amino acid sequence identity to hIL-1Raß. In addition, a portion of the DNA sequence of clone DNA102044 (Figure 16) (SEO ID NO:20) was found to coincide with the DNA sequence of EST AI014548 (Figure 4) (SEQ ID NO:8) and with the complement of the DNA sequence of EST Al343258 (Figure 17) (SEQ ID NO:23).

EST clone Al343258 was purchased from Research Genetics (Huntsville, AL) and the cDNA insert was obtained and sequenced in its entirety. The entire sequence of the clone

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AI343258, designated DNA114876, is shown in Figure 19 (SEQ ID NO:24). Clone DNA114876 contains a single open reading frame (ORF) with an apparent translation initiation site at nucleotide positions 73-75 and a stop codon at nucleotide positions 726-728 (Figure 19; SEQID NO:24), encoding a predicted polypeptide precursor (hIL-1Ra1V) (Fig. 19; SEQ ID NO:25) that is 218 amino acids long. In addition, the ORF contains an alternate translation initiation site at nucleotide positions 106-108. The predicted polypeptide (also designated hIL-1Ra1V) for translation initiated at the alternate start codon is 207 amino acids in length (lacking the first eleven residues at the N-terminus of the 218 amino acid polypeptide). It is believed that the predicted 218 amino acid and 207 amino acid polypeptides behave as mature sequences (without a presequence that is removed in posttranslational processing) in certain animal cells. It is also believed that other animal cells recognize and remove one or more signal peptide(s) extending from amino acid positions 1 to about 48 (a putative leader sequence in the 218 amino acid polypeptide) or from amino acid positions 12 to 36 (a putative leader sequence in the 207 amino acid polypeptide) in the amino acid sequence of Figure 19 (SEQ ID NO:25). As shown in Example 14 below, transiently transfected CHO host cells secrete unprocessed forms of hlL-1Ra1V and hlL-1Ra1L and a single processed form that results from the removal of a signal peptide extending from amino acid positions 1 to 45 in Figure 19 (SEQ ID NO:25) or the removal of a signal peptide extending from amino acid positions 1 to 34 of Figure 15 (SEQ ID NO:19). The processed form of hIL-1Ra1V and hIL-1Ra1L secreted by transiently transfected CHO host cells has the amino acid sequence of amino acid residues 35 to 207 of Figure 15 (SEQ ID NO:19) and amino acid residues 46 to 218 of Figure 19 (SEQ ID NO:25).

Clone DNA114876 (designated DNA114876-2534) was deposited with ATCC and was assigned ATCC deposit no. 203973. The full length hlL-1Ra1V protein shown in Figure 19 (SEQ ID NO:25) has an estimated molecular weight of about 24,124 and a pl of about 6.1.

Based on a sequence alignment analysis of the full length sequence (SEQ ID NO:25), hIL-1Ra1V shows significant amino acid sequence identity to hIL-1Raβ. hIL-1Ra1V is believed to be an allelic variant of hIL-1Ra1L.

## EXAMPLE 13

### IL-18 Receptor and IL-1Receptor Binding of hIL-1Ra1S

To facilitate the characterization of hll.-IRa1S, a PCR fragment encoding amino acid residues 39-167 in the ORF of clone DNA102044 (Figure 16; SEQ ID NO:21) was cloned into pCMVIFLAG (IBI Kodak, described in Pan et al., Science, 276: 111-113) as an in-frame fusion to a NH<sub>2</sub>-terminal preprotryps in leader sequence and FLAG tag encoded by the vector to form plasmid pCMVIFLAG-IL-IRa1S. Plasmid pCMVIFLAG-IL18R-ECD-Fc was obtained as described in Example 9 above.

Human embryonic kidney 293 cells were grown in high glucose DMEM (Genentech, Inc). The cells were seeded at 3-4 X10° per plate (100 mm) and co-transfected with pCMV1FLAG-IIL1Ra1S and pCMV1FLAG-IIL18R.ECD-Fc by means of calcium phosphate precipitation. The media were changed 12 hours post transfection. The resultant conditioned media (10 ml each) were harvested after a further 70-74 hour incubation, clarified by centrifugation, aliquoted and stored at -70°C. The receptor-Fc and ligand complex from 1.5 ml conditioned medium was

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immunoprecipitated with protein G-Sepharose, washed three times with buffer containing 50 mM Hepes, pH7.0, 150 mM NaCl, 1 mM EDTA, 1% NP-40, and a protease inhibitor cocktail (BMB) and resolved on a 10-20% SDS-PAGE gel. The bound ligand was identified by immunoblotting using anti-FLAG monoclonal antibody (BMB).

The immunoblotting results indicated that the secreted FLAGhIL-1Ra1S fusion protein bound to IL-18R ECD. These data show that hIL-1Ra1S could be an agonist or antagonist of IL-18R.

#### EXAMPLE 14

## hIL-1Ra1V, hIL-1Ra1L and hIL-1Ra3 Processing

cDNAs encoding full-length hIL-1Ra1V (amino acids 1-218 in the ORF of clone DNA114876 shown in Figure 19 (SEQ ID NO:25)), full length hIL-1Ra1L (amino acids 1-207 in the ORF of clone DNA102043 shown in Figure 15 (SEQ ID NO:19)), and full length hIL-1Ra3 (amino acids 1-155 in the ORF of clone DNA96787 shown in Figure 7 (SEQ ID NO:13)) were each cloned into a pRK7 expression vector as an in-frame fusion with a carboxy-terminal FLAG-tag sequence. In preparation for mammalian cell transient transfections, CHO DP12 cells were seeded at 4 x 106 cells per plate (100mm petri dish) in growth medium (PS20, 5% FBS, 1X GHT, 1X pen/strep, 1X L-glutamine) the day before transfection. On the day of transfection, cells were washed with PBS and fed with 10 ml serum-free transfection medium (PS20, 1X GHT). DNAlipid transfection mixtures were prepared by adding stepwise into eppendorf tubes (1) 400 µl transfection medium (PS20, 1X GHT); (2) 12 µg DNA; (3) 10 µg poly-lysine; and (4) 50 µl Dosper liposomal transfection reagent (Boehringer Mannheim). The DNA-lipid mixtures were incubated for 15 minutes at room temperature and then added dropwise to cell culture plates. Cells were incubated overnight at 37°C. On the day after transfection, cells were washed with PBS, fed with 10 ml serum-free production medium (PS24, 10 mg/L insulin, 1X trace elements, 1.4 mg/L lipid EtOH), and placed in a 32°C incubator. After 5 days, the culture media containing the expressed proteins were harvested and cleared by centrifugation. For peptide sequencing of each expressed protein, 5-10 ml of the conditioned medium containing the expressed protein was incubated with monoclonal anti-FLAG antibody (Boehringer Mannheim) coupled to agarose beads. The immunoprecipitated FLAG-tag proteins were extensively washed with 1% NP-40 buffer (125 mM NaCl, 1 mM EDTA and 50 mM Tris-HCl, pH 7.4). The immunoprecipitates were run on a SDS polyacrylamide gel, the separated polypeptides on the gel were transferred to a PVDF membrane, the PVDF membrane was stained with Coomassie blue, and the corresponding protein bands were excised from the membrane. The amino-terminal protein sequences were obtained by conventional methods.

The processed N-terminal sequence of both of the hIL-1Ra1L and hIL-1Ra1V polypeptides was determined to be VHTSPKVKN (SEQ ID NO:31). Approximately 50% of hIL-1Ra1L and hIL-1Ra1V material recovered from conditioned media exhibited the processed N-terminal sequence, indicating that the CHO host cells secreted a processed form corresponding to amino acid residues 35 to 207 in the amino acid sequence of Figure 15 (SEQ ID NO:19) and amino acid residues 46 to 218 in the amino acid sequence of Figure 19 (SEQ ID NO:25). The remaining 50% of the hIL-1Ra1L and hIL-1Ra1V material recovered from conditioned media exhibited an

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unprocessed N-terminus, indicating that the CHO host cells also secreted unprocessed forms

unprocessed N-terminus, indicating that the CHO host cells also secreted unprocessed forms of hLI-IRaIL and hIL-IRaIV corresponding to amino acid residues 1 to 207 in the amino acid sequence of Figure 15 (SEQ ID NO:19) and to amino acid residues 1 to 218 in the amino acid sequence of Figure 19 (SEQ ID NO:25), respectively.

The processed N-terminal sequence of both of the hIL-1Ra3 and mIL-1Ra3 polypeptides was determined to be VLSGALCFRM (SEQ ID NO.32). Approximately 100% of the hIL-1Ra3 and mIL-1Ra3 material recovered from conditioned media exhibited the processed N-terminal sequence, indicating that the CHO host cells secreted processed forms of hIL-1Ra3 and mIIL-1Ra3 that lack the N-terminal methionine and correspond to amino acid residues 2 to 155 in the amino acid sequence of Figure 7 (SEQ ID NO:13) and amino acid residues 2 to 155 in the amino acid sequence of Figure 9 (SEO ID NO:16), respectively.

## Deposit of Material

The following materials have been deposited with the American Type Culture Collection, 10801 University Blvd., Manassas, VA 20110-2209, USA (ATCC):

15	Material	ATCC Dep. No.	Deposit Date
	pSPORT1-based plasmid DNA92929-2534	203586	Jan. 12,1999
20	pCMV-1Flag-pcmv5 plasmid DNA96786-2534	203587	Jan. 12, 1999
	pT7T3D-Pac plasmid DNA85066-2534	203588	Jan. 12, 1999
25	pINCY-based plasmid DNA96787-2534	203589	Jan. 12, 1999
30	pT7T3D-Pac plasmid DNA92505-2534	203590	Jan. 12, 1999
30	pRK7-based plasmid DNA102043-2534	203846	March 16, 1999
35	pRK7-based plasmid DNA102044-2534	203855	March 16, 1999
	pRK7-based plasmid DNA114876-2534	203973	April 27, 1999

These deposits were made under the provisions of the Budapest Treaty on the International Recognition of the Deposit of Microorganisms for the Purpose of Patent Procedure and the Regulations thereunder (Budapest Treaty). This assures maintenance of viable cultures of the deposits for 30 years from the date of deposit. The deposits will be made available by ATCC under the terms of the Budapest Treaty, and subject to an agreement between Genentech, Inc. and ATCC, which assures permanent and unrestricted availability of the progeny of the cultures of the deposits to the public upon issuance of the pertinent U.S. patent or upon laying open to the public of any U.S. or foreign patent application, whichever comes first, and assures availability of the progeny to one determined by the U.S. Commissioner of Patents and

Trademarks to be entitled thereto according to 35 USC  $\S122$  and the Commissioner's rules pursuant thereto (including 37 CFR  $\S1.14$  with particular reference to 886 OG 638).

The assignee of the present application has agreed that if a culture of the materials on deposit should die or be lost or destroyed when cultivated under suitable conditions, the materials will be promptly replaced on notification with another of the same. Availability of the deposited material is not to be construed as a license to practice the invention in contravention of the rights granted under the authority of any government in accordance with its patent laws.

The foregoing written specification is considered to be sufficient to enable one skilled in the art to practice the invention. The present invention is not to be limited in scope by the constructs deposited, since the deposited embodiment is intended as a single illustration of certain aspects of the invention and any constructs that are functionally equivalent are within the scope of this invention. The deposit of material herein does not constitute an admission that the written description herein contained is inadequate to enable the practice of any aspect of the invention, including the best mode thereof, nor is it to be construed as limiting the scope of the claims to the specific illustrations that it represents. Indeed, various modifications of the invention in addition to those shown and described herein will become apparent to those skilled in the art from the foregoing description and fall within the scope of the appended claims.

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## Table 2A

PRO XXXXXXXXXXXXXX (Length = 15 amino acids) Comparison Protein XXXXYYYYYYY (Length = 12 amino acids)

% amino acid sequence identity =

(the number of identically matching amino acid residues between the two polypeptide sequences as determined by ALIGN-2) divided by (the total number of amino acid residues of the PRO polypeptide) =

5 divided by 15 = 33.3%

## Table 2B

% amino acid sequence identity =

(the number of identically matching amino acid residues between the two polypeptide sequences as determined by ALIGN-2) divided by (the total number of amino acid residues of the PRO polypeptide) =

5 divided by 10 = 50%

## Table 2C

PRO-DNA NNNNNNNNNNNNN (Length = 14 nucleotides)
Comparison DNA NNNNNLLLLLLLLLL (Length = 16 nucleotides)

% nucleic acid sequence identity =

(the number of identically matching nucleotides between the two nucleic acid sequences as determined by ALIGN-2) divided by (the total number of nucleotides of the PRO-DNA nucleic acid sequence) =

35 6 divided by 14 = 42.9%

# Table 2D

PRO-DNA NNNNNNNNNN (Length = 12 nucleotides)
Comparison DNA NNNLLLVV (Length = 9 nucleotides)

% nucleic acid sequence identity =

(the number of identically matching nucleotides between the two nucleic acid sequences as determined by ALIGN-2) divided by (the total number of nucleotides of the PRO-DNA nucleic acid sequence) =

4 5 4 divided by 12 = 33.3%

}; 40

## Table 3A

```
* C-C increased from 12 to 15
           * Z is average of EQ
           * B is average of ND
           * match with stop is M; stop-stop = 0; J (joker) match = 0
          #define M
                                  /* value of a match with a stop */
  10
          int
                   day[26][26] = {
                ABCDEF GHIJK LM NOPQR STUV W X Y Z */
                    { 2, 0,-2, 0, 0,-4, 1,-1,-1, 0,-1,-2,-1, 0,_M, 1, 0,-2, 1, 1, 0, 0,-6, 0,-3, 0},
                    { 0, 3,-4, 3, 2,-5, 0, 1,-2, 0, 0,-3,-2, 2, M,-1, 1, 0, 0, 0, 0,-2,-5, 0,-3, 1},
          /* B */
                    {-2,-4,15,-5,-5,-4,-3,-3,-2, 0,-5,-6,-5,-4, M,-3,-5,-4, 0,-2, 0,-2,-8, 0, 0,-5},
   15
          /* C */
          /* D */
                    { 0, 3,-5, 4, 3,-6, 1, 1,-2, 0, 0,-4,-3, 2, M,-1, 2,-1, 0, 0, 0,-2,-7, 0,-4, 2},
          /* E */
                    { 0, 2, -5, 3, 4, -5, 0, 1, -2, 0, 0, -3, -2, 1, M, -1, 2, -1, 0, 0, 0, -2, -7, 0, -4, 3},
                    {-4,-5,-4,-6,-5, 9,-5,-2, 1, 0,-5, 2, 0,-4, M,-5,-5,-4,-3,-3, 0,-1, 0, 0, 7,-5},
          /* F */
                    {1,0,-3,1,0,-5,5,-2,-3,0,-2,-4,-3,0,M,-1,-1,-3,1,0,0,-1,-7,0,-5,0},
          /* G */
                    {-1, 1,-3, 1, 1,-2,-2, 6,-2, 0, 0,-2,-2, 2, M, 0, 3, 2,-1,-1, 0,-2,-3, 0, 0, 2},
   20
          /* H */
                    {-1,-2,-2,-2,-1,-3,-2, 5, 0,-2, 2, 2,-2, M,-2,-2,-1, 0, 0, 4,-5, 0,-1,-2},
          /* I */
                    /* J */
                    {-1, 0,-5, 0, 0,-5,-2, 0,-2, 0, 5,-3, 0, 1, M,-1, 1, 3, 0, 0, 0,-2,-3, 0,-4, 0},
          /* K */
                    {-2,-3,-6,-4,-3, 2,-4,-2, 2, 0,-3, 6, 4,-3, M,-3,-2,-3,-3,-1, 0, 2,-2, 0,-1,-2},
          /* L */
                    {-1,-2,-5,-3,-2, 0,-3,-2, 2, 0, 0, 4, 6,-2,_M,-2,-1, 0,-2,-1, 0, 2,-4, 0,-2,-1},
          /* M */
                    { 0, 2, 4, 2, 1, 4, 0, 2, -2, 0, 1, -3, -2, 2, M, -1, 1, 0, 1, 0, 0, -2, -4, 0, -2, 1},
          /* N */
                    /* O */
43
                    { 1,-1,-3,-1,-1,-5,-1, 0,-2, 0,-1,-3,-2,-1, M, 6, 0, 0, 1, 0, 0,-1,-6, 0,-5, 0},
          /* P */
1.7
                    { 0, 1,-5, 2, 2,-5,-1, 3,-2, 0, 1,-2,-1, 1, M, 0, 4, 1,-1,-1, 0,-2,-5, 0,-4, 3},
          /* O */
111 30
                    {-2, 0,-4,-1,-1,-4,-3, 2,-2, 0, 3,-3, 0, 0, M, 0, 1, 6, 0,-1, 0,-2, 2, 0,-4, 0},
          /* R */
                    U
          /* S */
          /* T */
          /* U */
                    \{0,-2,-2,-2,-1,-1,-2,4,0,-2,2,2,2,-M,-1,-2,-2,-1,0,0,4,-6,0,-2,-2\},
          /* V */
   35
                     {-6,-5,-8,-7,-7, 0,-7,-3,-5, 0,-3,-2,-4,-4, M,-6,-5, 2,-2,-5, 0,-6,17, 0, 0,-6},
          /* W */
                    14
          /* X */
                    {-3,-3, 0,-4,-4, 7,-5, 0,-1, 0,-4,-1,-2,-2, M,-5,-4,-4,-3,-3, 0,-2, 0, 0.10,-4}.
150
          /* Y */
/* Z */
                    {0, 1,-5, 2, 3,-5, 0, 2,-2, 0, 0,-2,-1, 1, M, 0, 3, 0, 0, 0, 0, -2,-6, 0,-4, 4}
```

char

55 char

## Table 3B

```
#include < stdio.h >
           #include < ctype.h >
                                        16
                                                  /* max jumps in a diag */
           #define MAXJMP
                                        24
                                                  /* don't continue to penalize gaps larger than this */
           #define MAXGAP
           #define JMPS
                                        1024
                                                  /* max jmps in an path */
   10
           #define MX
                                        4
                                                  /* save if there's at least MX-1 bases since last jmp */
           #define DMAT
                                        3
                                                  /* value of matching bases */
           #define DMIS
                                        0
                                                  /* penalty for mismatched bases */
           #define DINSO
                                        8
                                                  /* penalty for a gap */
   15
           #define DINS1
                                                  /* penalty per base */
                                        1
           #define PINSO
                                        8
                                                  /* penalty for a gap */
           #define PINS1
                                                  /* penalty per residue */
           struct jmp {
   20
                                        nfMAXJMP1:
                                                           /* size of jmp (neg for dely) */
                                        x[MAXJMP];
                                                           /* base no. of imp in seq x */
                     unsigned short
                                                           /* limits seq to 2^16 -1 */
10
10
10
10
10
10
10
           struct diag {
                                                           /* score at last imp */
                                        score:
                     long
                                        offset;
                                                           /* offset of prev block */
                                                           /* current imp index */
                     short
                                         ijmp;
                                                           /* list of jmps */
                     struct jmp
                                        jp;
30
           struct path {
13
                     int
                                                  /* number of leading spaces */
                     short
                               n[JMPS]; /* size of imp (gap) */
                               x[JMPS]; /* loc of jmp (last elem before gap) */
N 35
            };
13
            char
                               *ofile:
                                                            /* output file name */
ıØ
            char
                               *namex[2];
                                                            /* seg names: getsegs() */
            char
                               *prog;
                                                            /* prog name for err msgs */
140
            char
                               *seqx[2];
                                                           /* seqs: getseqs() */
            int
                               dmax;
                                                           /* best diag: nw() */
            int
                               dmax0:
                                                            /* final diag */
            int
                                                            /* set if dna: main() */
                               dna;
            int
                                                            /* set if penalizing end gaps */
                               endgaps;
   4.5
                                                            /* total gaps in seqs */
            int
                               gapx, gapy;
            int
                               len0, len1;
                                                            /* seq lens */
                                                            /* total size of gaps */
            int
                               ngapx, ngapy;
                                                            /* max score: nw() */
            int
                               smax:
                               *xbm;
                                                            /* bitmap for matching */
            int
   50
            long
                               offset;
                                                           /* current offset in imp file */
                                                            /* holds diagonals */
            struct
                     diag
                               *dx:
            struct
                     path
                               pp[2]:
                                                           /* holds path for seqs */
```

\*calloc(), \*malloc(), \*index(), \*strcpy();

\*getseq(), \*g calloc();

# Table 3C

```
/* Needleman-Wunsch alignment program
             * usage: progs file1 file2
      5
             * where file1 and file2 are two dna or two protein sequences.
             * The sequences can be in upper- or lower-case an may contain ambiguity
             * Any lines beginning with ';', '>' or '<' are ignored
             * Max file length is 65535 (limited by unsigned short x in the jmp struct)
                A sequence with 1/3 or more of its elements ACGTU is assumed to be DNA
    10
             * Output is in the file "align.out"
             * The program may create a tmp file in /tmp to hold info about traceback.
             * Original version developed under BSD 4.3 on a vax 8650
    15
            #include "nw.h"
            #include "day.h"
            static
                     dbval[26] = {
                     1,14,2,13,0,0,4,11,0,0,12,0,3,15,0,0,0,5,6,8,8,7,9,0,10.0
    20
            };
            static
                     _pbval[26] = {
                     1. 21(1 < <('D'-'A'))1(1 < <('N'-'A')), 4, 8, 16, 32, 64.
                     128, 256, 0xFFFFFFF, 1 < < 10, 1 < < 11, 1 < < 12, 1 < < 13, 1 < < 14,
    25
                     1<<15, 1<<16, 1<<17, 1<<18, 1<<19, 1<<20, 1<<21, 1<<22,
                     1< <23, 1< <24, 1< <25[(1<<('E'-'A'))](1<<('Q'-'A'))
            };
                                                                                                                           main
            main(ac, av)
m 30
                              ac:
                     char
                              *av[]:
                     prog = av[0];
                     if (ac! = 3) {
    35
                               fprintf(stderr, "usage: %s file1 file2\n", prog);
                               fprintf(stderr, "where file1 and file2 are two dna or two protein sequences.\n");
                               fprintf(stderr, "The sequences can be in upper- or lower-case\n");
                               fprintf(stderr, "Any lines beginning with ';' or ' < ' are ignored\n");
                               fprintf(stderr, "Output is in the file \"align.out\"\n");
    40
                              exit(1):
                     namex[0] = av[1];
                     namex[1] = av[2];
                     seqx[0] = getseq(namex[0], &len0);
    45
                     seqx[1] = getseq(namex[1], &len1);
                     xbm = (dna)? dbval : pbval;
                     endgaps \approx 0:
                                                           /* 1 to penalize endgaps */
                     ofile = "align.out";
                                                          /* output file */
    50
                                        /* fill in the matrix, get the possible imps */
                     nw();
                     readimps();
                                        /* get the actual imps */
                     print();
                                       /* print stats, alignment */
    55
                     cleanup(0):
                                       /* unlink any tmp files */
```

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## Table 3D

```
/* do the alignment, return best score: main()
             * dna: values in Fitch and Smith, PNAS, 80, 1382-1386, 1983
             * pro: PAM 250 values
             * When scores are equal, we prefer mismatches to any gap, prefer
             * a new gap to extending an ongoing gap, and prefer a gap in seqx
             * to a gap in seq y.
             */
                                                                                                                               nw
            nw()
            {
                      char
                                         *px, *py;
                                                                      /* seqs and ptrs */
                                         *ndely, *dely;
                                                            /* keep track of dely */
                      int
                      int
                                         ndelx, delx;
                                                            /* keep track of delx */
                                                            /* for swapping row0, row1 */
                      int
                                         *tmp;
    15
                      int
                                         mis;
                                                            /* score for each type */
                      int
                                         ins0, ins1:
                                                            /* insertion penalties */
                      register
                                         id;
                                                            /* diagonal index */
                                                            /* jmp index */
                      register
                                         ij;
                      register
                                         *col0, *col1;
                                                            /* score for curr, last row */
    20
                                                            /* index into segs */
                      register
                                         xx, yy;
                      dx = (struct diag *)g_calloc("to get diags", len0+len1+1, sizeof(struct diag));
00
                      ndely = (int *)g calloc("to get ndely", len1+1, sizeof(int));
    25
                      dely = (int *)g_calloc("to get dely", len1+1, sizeof(int));
                      col0 = (int *)g calloc("to get col0", len1 +1, sizeof(int));
m
                      col1 = (int *)g calloc("to get col1", len1+1, sizeof(int));
                      ins0 = (dna)? DINS0 : PINS0;
U
                      ins1 = (dna)? DINS1 : PINS1;
13
    30
D)
                      smax = -10000:
                      if (endgaps) {
                                for (col0[0] = dely[0] = -ins0, yy = 1; yy < = len1; yy + +) {
27
                                         col0[yy] = dely[yy] = col0[yy-1] - ins1;
n
    35
                                         ndely[yy] = yy;
10
902
                               col0[0] = 0;
                                                   /* Waterman Bull Math Biol 84 */
                       else
    40
                                for (yy = 1; yy \le len1; yy++)
                                         dely[yy] = -ins0;
                       /* fill in match matrix
                       for (px = seqx[0], xx = 1; xx <= len0; px++, xx++) {
    45
                                /* initialize first entry in col
                                 */
                                if (endgaps) {
                                          if (xx == 1)
    50
                                                   col1[0] = delx = -(ins0 + ins1);
                                                   col1[0] = delx = col0[0] - ins1;
                                          ndelx = xx
     55
                                else {
                                          col1[0] = 0;
                                          delx = -ins0;
                                          ndelx = 0:
                                }
```

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## Table 3E

```
...nw
for (py = seqx[1], yy = 1; yy < = len1; py++, yy++) {
         mis = col0[yy-1];
         if (dna)
                  mis += (xbm[*px-'A']&xbm[*py-'A'])? DMAT : DMIS;
         else
                  mis += _day[*px-'A'][*py-'A'];
         /* update penalty for del in x seq;
         * favor new del over ongong del
         * ignore MAXGAP if weighting endgaps
         if (endgaps | | ndely[yy] < MAXGAP) {
                  if (col0[yy] - ins0 > = dely[yy]) {
                           dely[yy] = col0[yy] - (ins0 + ins1);
                           ndely[yy] = 1;
                  } else {
                           dely[yy] -= ins1;
                           ndely[yy]++;
         } else {
                  if (col0[yy] - (ins0 + ins1) > = dely[yy]) {
                           dely[yy] = col0[yy] - (ins0+ins1);
                           ndelv[yy] = 1;
                  } else
                           ndely[yy]++;
         }
         /* update penalty for del in y seq;
         * favor new del over ongong del
         if (endgaps | | ndelx < MAXGAP) {
                  if (col1[yy-1] - ins0 > = delx) {
                           delx = col1[vv-1] - (ins0+ins1);
                           ndelx == 1:
                  } else {
                           delx -= insl:
                           ndelx++;
         } else {
                  if (coll[yy-1] - (ins0+ins1) > = delx) {
                           delx = col1[vy-1] - (ins0+ins1);
                           ndelx = 1:
                  } else
                           ndelx++;
         /* pick the maximum score; we're favoring
         * mis over any del and delx over dely
         */
                                Table 3F
                                                                                         ...nw
         id = xx - yy + len1 - 1;
         if (mis > = delx && mis > = delv[vv])
                  coll{yy] = mis;
         else if (delx > = dely[yy]) {
                  coll[yy] = delx;
                  ij = dx[id].ijmp;
                  if (dx[id], ip.n[0] && (!dna | | (ndelx > = MAXJMP))
                  && xx > dx[id].ip.x[ij]+MX) \mid | mis > dx[id].score+DINS0)) {
                           dx[id].ijmp++;
                           if (++ij > = MAXJMP) {
```

-102-

}

```
writejmps(id);
                                                                    ij = dx[id].ijmp = 0;
                                                                    dx[id].offset = offset;
                                                                    offset += sizeof(struct jmp) + sizeof(offset);
      5
                                                           }
                                                 dx[id].jp.n[ij] = ndelx;
                                                 dx[id].jp.x[ij] = xx;
                                                 dxfid1.score = delx;
    10
                                        else {
                                                 col1[yy] = dely[yy];
                                                 ij = dx[id].ijmp;
    15
                      if (dx[id].jp.n[0] && (!dna \mid | (ndely[yy]) > = MAXJMP
                                                 && xx > dx[id].jp.x[ij]+MX) \mid \mid mis > dx[id].score+DINS0)) {
                                                           dx[id].ijmp++;
                                                           if (++ij > = MAXJMP) {
                                                                    writejmps(id);
    20
                                                                    ij = dx[id].ijmp = 0;
                                                                    dx[id].offset = offset;
                                                                    offset += sizeof(struct imp) + sizeof(offset);
                                                           3
25
                                                 dx[id].jp.n[ij] = -ndely[yy];
                                                 dx[id].jp.x[ij] = xx;
                                                 dx[id].score = dely[yy];
                                        if (xx = = len0 && yy < len1) {
    30
                                                 /* last col
                                                  */
                                                 if (endgaps)
                                                           coll[yy] -= ins0 + ins1*(len1-yy);
if (coll[yy] > smax) {
    35
                                                           smax = col1[yy];
                                                           dmax = id;
    40
                               if (endgaps && xx < len0)
                                        col1[yy-1] -= ins0 + ins1*(len0-xx);
                               if (coll[yy-1] > smax) {
                                         smax = col1[yy-1];
                                        dmax = id:
     45
                               tmp = col0; col0 = col1; col1 = tmp;
                      (void) free((char *)ndely);
                      (void) free((char *)dely);
     50
                      (void) free((char *)col0);
                      (void) free((char *)col1);
```

# Table 3G

```
/4
             * print() -- only routine visible outside this module
      5
             * getmat() -- trace back best path, count matches: print()
             * pr align() -- print alignment of described in array p[]: print()
             * dumpblock() -- dump a block of lines with numbers, stars: pr_align()
    10
             * nums() -- put out a number line: dumpblock()
             * putline() -- put out a line (name, [num], seq, [num]): dumpblock()
             * stars() - -put a line of stars: dumpblock()
             * stripname() - strip any path and prefix from a sequame
    15
            #include "nw.h*
            #define SPC
            #define P LINE 256
                                         /* maximum output line */
    20
            #define P SPC
                               3
                                         /* space between name or num and seq */
            extern
                      _day[26][26];
                                         /* set output line length */
            int
                      olen:
口
道 25
切
切
            FILE
                      *fx:
                                         /* output file */
            print()
                                                                                                                               print
            {
S
                      int
                                lx, ly, firstgap, lastgap;
                                                            /* overlap */
    30
                      if ((fx = fopen(ofile, "w")) == 0) {
                                fprintf(stderr, "%s: can't write %s\n", prog, ofile);
(0)
                                cleanup(1);
                      fprintf(fx, " < first sequence: %s (length = %d)\n", namex[0], len0);
                      fprintf(fx, "<second sequence: %s (length = %d)\n", namex[1], len1);
    35
                      olen = 60;
1 de
                      lx = len0;
100
                      ly = len1;
1 40
                      firstgap = lastgap = 0;
                      if (dmax < len1 - 1) {
                                                  /* leading gap in x */
                                pp[0].spc = firstgap = len1 - dmax - 1;
                                ly -= pp[0].spc;
                      else if (dmax > len1 - 1) { /* leading gap in y */
    45
                                pp[1].spc = firstgap = dmax - (len1 - 1);
                                lx -= pp[1].spc;
                      if (dmax0 < len0 - 1) {
                                                  /* trailing gap in x */
                                lastgap = len0 - dmax0 - 1;
    50
                                lx -= lastgap;
                      else if (dmax0 > len0 - 1) { /* trailing gap in y */
                                lastgap = dmax0 - (len0 - 1);
                                ly -= lastgap;
    55
                      getmat(lx, ly, firstgap, lastgap);
                      pr align();
             }
```

# Table 3H

```
* trace back the best path, count matches
     5
                                                                                                                        getmat
            getmat(lx, ly, firstgap, lastgap)
                     int
                              lx, ly;
                                                          /* "core" (minus endgaps) */
                                                          /* leading trailing overlap */
                              firstgap, lastgap;
   10
                                       nm, i0, i1, siz0, siz1;
                     int
                     char
                                       outx[32];
                     double
                                       pct;
                                       n0, n1;
                     register
                     register char
                                        *p0, *p1;
   15
                     /* get total matches, score
                     i0 = i1 = siz0 = siz1 = 0;
                     p0 = seqx[0] + pp[1].spc;
   20
                     p1 = seqx[1] + pp[0].spc;
                     n0 = pp[1].spc + 1;
                     n1 = pp[0].spc + 1;
10 25 m 40 m 30 m
                     nm = 0;
                     while ( *p0 && *p1 ) {
                              if (siz0) {
                                        p1++;
                                        n1++;
                                        siz0--:
                               else if (siz1) {
                                        p0++;
. 13
                                        n0++;
                                        siz1--;
    35
                               else {
14
                                        if (xbm[*p0-'A']&xbm[*p1-'A'])
90
                                                 nm++;
                                        if(n0++==pp[0].x[i0])
11 40
                                                siz0 = pp[0].n[i0++];
                                        if (n1++==pp[1].x[i1])
                                                 siz1 = pp[1].n[i1++];
                                        p0++;
                                        p1++;
    45
                               }
                     /* pct homology:
                      * if penalizing endgaps, base is the shorter seq
    50
                      * else, knock off overhangs and take shorter core
                      */
                     if (endgaps)
                               1x = (1en0 < 1en1)? len0 : len1;
    55
                               1x = (1x < 1y)? 1x : 1y;
                     pct = 100.*(double)nm/(double)lx;
                      fprintf(fx, "\n");
                     fprintf(fx, " < %d match%s in an overlap of %d: %.2f percent similarity\n",
                               nm, (nm == 1)? "" : "es", lx, pct);
```

PCT/US99/30720 WO 00/39297

#### Table 3I

```
...getmat
                    fprintf(fx, " < gaps in first sequence: %d", gapx);
                    if (gapx) {
    5
                              (void) sprintf(outx, " (%d %s%s)",
                                       ngapx, (dna)? "base": "residue", (ngapx == 1)? "": "s");
                              forintf(fx, "%s", outx);
                    fprintf(fx, ", gaps in second sequence: %d", gapy);
  10
                    if (gapy) {
                              (void) sprintf(outx, " (%d %s%s)",
                                       ngapy, (dna)? "base": "residue", (ngapy == 1)? "": "s");
                              fprintf(fx, "%s", outx);
   15
                    if (dna)
                              fprintf(fx,
                              "\n < score: %d (match = %d, mismatch = %d, gap penalty = %d + %d per base)\n",
                              smax, DMAT, DMIS, DINSO, DINS1);
                    else
   20
                              fprintf(fx.
                              "\n < score: %d (Dayhoff PAM 250 matrix, gap penalty = %d + %d per residue)\n",
                              smax, PINS0, PINS1);
                     if (endgaps)
                              fprintf(fx,
                              "<endgaps penalized. left endgap: %d %s%s, right endgap: %d %s%s\n",
   25
                              firstgap, (dna)? "base": "residue", (firstgap = = 1)? "": "s",
                              lastgap, (dna)? "base" : "residue", (lastgap = = 1)? "" : "s");
                     else
                              fprintf(fx, " < endgaps not penalized\n");
    30
                                                 /* matches in core -- for checking */
            static
                              nm:
                                                 /* lengths of stripped file names */
            static
                              lmax:
                                                 /* jmp index for a path */
            static
                               ii[2];
                                                 /* number at start of current line */
   35
            static
                              nc[2];
                                                 /* current elem number -- for gapping */
            static
                               ni[2];
            static
                               siz[2];
                                                 /* ptr to current element */
            static char
                               *ps[2];
                                                 /* ptr to next output char slot */
            static char
                               *po[2];
NJ 40
                               out[2][P_LINE]; /* output line */
            static char
                               star[P_LINE];
                                                 /* set by stars() */
            static char
             * print alignment of described in struct path pp[]
    45
            static
                                                                                                                       pr align
            pr_align()
                                                  /* char count */
                      int
                                        nn:
    50
                      int
                                        more:
                      register
                                         i:
                      for (i = 0, lmax = 0; i < 2; i++)
                               nn = stripname(namex[i]);
                               if (nn > lmax)
                                         lmax = nn;
                               nc[i] = 1;
                               ni[i] = 1;
    60
                               siz[i] = ij[i] = 0;
                               ps[i] = seqx[i];
                               po[i] = out[i];
```

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### Table 3J

```
...pr align
                      for (nn = nm = 0, more = 1; more;)
                               for (i = more = 0; i < 2; i++) {
      5
                                         * do we have more of this sequence?
                                         */
                                        if (!*psfil)
                                                 continue;
    10
                                        more++;
                                        if (pp[i].spc) { /* leading space */
                                                  *po[i]++ = ' ';
                                                 pp[i].spc--;
    15
                                        else if (siz[i]) { /* in a gap */
                                                  *po[i]++ = '-';
                                                  siz[i]--;
    20
                                         else {
                                                          /* we're putting a seq element
                                                  po[i] = ps[i];
if (islower(*ps[i]))
     25
                                                          *ps[i] = toupper(*ps[i]);
                                                  ps[i]++;
    30
                                                  * are we at next gap for this seq?
                                                  if (ni[i] == pp[i].x[ij[i]]) \{
· DRI GON
                                                           * we need to merge all gaps
                                                           * at this location
     35
                                                           siz[i] = pp[i].n[ij[i]++];
                                                           while (ni[i] = pp[i].x[ij[i]])
                                                                    siz[i] += pp[i].n[ij[i]++];
     40
                                                  nifil++;
                                if (+ +nn == olen | | !more && nn) {
     45
                                         dumpblock();
                                         for (i = 0; i < 2; i++)
                                                 po[i] = out[i];
                                         nn = 0;
                                }
     50
                       }
              * dump a block of lines, including numbers, stars: pr align()
     55
              */
             static
                                                                                                                  dumpblock
             dumpblock()
                       register i;
     60
                       for (i = 0; i < 2; i++)
                                *po[i]-- = '\0';
```

# Table 3K

...dumpblock

```
(void) putc('\n', fx);
      5
                      for (i = 0; i < 2; i++)
                               if (*out[i] && (*out[i] != ' | | *(po[i]) != ' ')) {
                                         if (i = 0)
                                                   nums(i):
                                         if (i == 0 && *out[1])
    10
                                         nutline(i);
                                         if (i == 0 \&\& *out[1])
                                                   fprintf(fx, star);
                                         if (i == 1)
    15
                                                   nums(i):
                               }
                      }
            }
    20
             * put out a number line: dumpblock()
            static
nums
            nums(ix)
    25
                                         /* index in out[] holding seq line */
                                ix;
                                          nline[P LINE];
                      char
                      register
                                          i, j;
                      register char
                                          *pn, *px, *py;
(7) 30
                      for (pn = nline, i = 0; i < lmax + P_SPC; i++, pn++)
(7)
                                *pn = ' ':
                      for (i = nc[ix], py = out[ix]; *py; py++, pn++) {
    if (*py == ' ' || *py == '-')
    *pn = ' ';
    35
                                else {
900
                                          if (i\%10 == 0 \mid | (i == 1 \&\& nc[ix] != 1)) 
                                                   j = (i < 0)? -i : i;
                                                    for (px = pn; j; j /= 10, px--)
                                                              *px = j\%10 + '0';
    40
                                                    if (i < 0)
                                                              *px = '-';
                                          else
     45
                                                    *pn = ' ';
                                          i++;
                                }
                       *pn = '\0';
     50
                       nc[ix] = i;
                       for (pn = nline; *pn; pn++)
                                (void) putc(*pn, fx);
                       (void) putc('\n', fx);
             }
     55
              * put out a line (name, [num], seq, [num]): dumpblock()
             static
                                                                                                                              putline
     60
             putline(ix)
                       int
                                 ix;
             {
```

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\*px = '\0';

55

# Table 3L

```
...putline
                    int
     5
                    register char
                                      *px:
                    for (px = namex[ix], i = 0; *px && *px != ':'; px++, i++)
                             (void) putc(*px, fx);
                    for (; i < lmax + P_SPC; i++)
   10
                             (void) putc(' ', fx);
                    /* these count from 1:
                     * ni[] is current element (from 1)
                     * ncll is number at start of current line
   15
                    for (px = out[ix]; *px; px++)
                             (void) putc(*px&0x7F, fx);
                    (void) putc('\n', fx);
           }
   20
            * put a line of stars (seqs always in out[0], out[1]): dumpblock()
25
0
0
0
0
0
0
0
0
           static
                                                                                                                    stars
           stars()
                    register char
                                      *p0, *p1, cx, *px;
                    return:
                    px = star;
    35
                    for (i = lmax + P SPC; i; i--)
14
                             *px + + = ' ';
J
                    for (p0 = out[0], p1 = out[1]; *p0 && *p1; p0++, p1++) {
if (isalpha(*p0) && isalpha(*p1)) {
    40
                                       if (xbm[*p0-'A']&xbm[*p1-'A']) {
                                               cx = '*';
                                               nm++:
                                      élse if (!dna && _day[*p0-'A'][*p1-'A'] > 0)

cx = '...';
    45
                                       else
                                               cx = ' ':
                              }
    50
                              else
                                       cx = ' ':
                              *px++=cx;
                     *px + + = '\n';
```

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#### Table 3M

```
* strip path or prefix from pn, return len: pr align()
        static
                                                                                                                 stripname
        stripname(pn)
                                    /* file name (may be path) */
                 register char
                                    *px, *py;
10
                 pv = 0:
                 for (px = pn; *px; px++)
                          if (*px == '/')
                                   py = px + 1;
15
                 if (py)
                          (void) strcpy(pn, py);
                 return(strlen(pn));
        }
20
                                                            Table 3N
        * cleanup() -- cleanup any tmp file
         * getseq() - read in seq, set dna, len, maxlen
25
         * g calloc() -- calloc() with error checkin
         * readjmps() -- get the good jmps, from tmp file if necessary
         * write imps() - write a filled array of imps to a tmp file: nw()
        #include "nw.h"
30
        #include < sys/file.h>
                  *iname = "/tmp/homgXXXXXX";
                                                                 /* tmp file for jmps */
        char
        FILE
35
        int
                 cleanup();
                                                                 /* cleanup tmp file */
        long
                  lseek();
         * remove any tmp file if we blow
40
                                                                                                                    cleanup
        cleanup(i)
                           i;
        {
                  if (fj)
45
                           (void) unlink(jname);
                  exit(i);
        }
50
         * read, return ptr to seq, set dna, len, maxien
         * skip lines starting with ';', '<', or '>'
         * seq in upper or lower case
        char
                                                                                                                       getseq
55
        getseq(file, len)
                           *file:
                                    /* file name */
                  char
                  int
                           *len:
                                    /* seg len */
         {
                  char
                                     line[1024], *pseq;
 60
                  register char
                                     *px, *py;
                  int
                                     natge, tlen;
                  FILE
                                     *fp;
                  if ((fp = fopen(file, "r")) == 0) {
```

-110-

```
fprintf(stderr, "%s: can't read %s\n", prog, file);
                        exit(1);
                tlen = natgc = 0;
 5
                while (fgets(line, 1024, fp)) {
                        if (*line == ';' | | *line == '<' | | *line == '>')
                                 continue;
                        for (px = line; *px != '\n'; px++)
                                 if (isupper(*px) | | islower(*px))
10
                                          tien++;
                if ((pseq = malloc((unsigned)(tlen+6))) == 0) {
                        fprintf(stderr, "%s: malloc() failed to get %d bytes for %s\n", prog, tlen+6, file);
15
                pseq[0] = pseq[1] = pseq[2] = pseq[3] = '\0';
                                                       Table 3O
                                                                                                         ...getseq
20
                py = pseq + 4;
                *len = tlen;
                rewind(fp);
                while (fgets(line, 1024, fp)) {
                         if (*line == ';' || *line == '<' || *line == '>')
25
                                 continue;
                         for (px = line; *px != '\n'; px++) {
                                  if (isupper(*px))
                                          *py++ = *px:
30
                                  else if (islower(*px))
                                          *py++ = toupper(*px);
                                  if (index("ATGCU",*(py-1)))
                                          natgc++;
                         }
35
                 *py++ = '\0';
                 *py = '\0';
                (void) fclose(fp);
                dna = natgc > (tlen/3);
40
                return(pseq+4);
        }
        char
                                                                                                          g_calloc
        g calloc(msg, nx, sz)
45
                                           /* program, calling routine */
                char
                         *msg:
                                           /* number and size of elements */
                 int
                         nx, sz;
                                  *px, *calloc();
                 char
50
                 if ((px = calloc((unsigned)nx, (unsigned)sz)) == 0) {
                         if (*msg) {
                                  exit(1);
55
                 }
                 return(px);
        }
 60
         * get final jmps from dx[] or tmp file, set pp[], reset dmax: main()
                                                                                                        readimps
        readjmps()
                 int
                                  fd = -1;
                                                            -111-
```

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```
siz. i0. i1:
                inf
                register i, j, xx;
                if (fj) {
 5
                         (void) fclose(fi);
                         if ((fd = open(jname, O RDONLY, 0)) < 0) {
                                  fprintf(stderr, "%s: can't open() %s\n", prog, jname);
                                  cleanup(1);
10
                 for (i = i0 = i1 = 0, dmax0 = dmax, xx = len0; ; i++) {
                         while (1) {
                                  for (j = dx[dmax].ijmp; j > = 0 && dx[dmax].jp.x[j] > = xx; j--)
15
                                                         Table 3P
                                                                                                          ...readimps
                                   if (j < 0 && dx[dmax].offset && fj) {
                                            (void) lseek(fd, dx[dmax].offset, 0);
20
                                            (void) read(fd, (char *)&dx[dmax].jp, sizeof(struct jmp));
                                            (void) read(fd, (char *)&dx[dmax].offset, sizeof(dx[dmax].offset));
                                            dx[dmax].ijmp = MAXJMP-1;
                                   }
                                   else
25
                                            break:
                          if (i > = JMPS) {
                                   fprintf(stderr, "%s: too many gaps in alignment\n", prog);
                                   cleanup(1);
30
                          if (i > = 0) {
                                   siz = dx[dmax].jp.n[j];
                                   xx = dx[dmax].jp.x[j];
                                   dmax += siz;
                                                              /* gap in second seq */
35
                                   if (siz < 0) {
                                            pp[1].n[i1] = -siz;
                                            xx += siz;
                                            /* id = xx - yy + len1 - 1
40
                                            pp[1].x[i1] = xx - dmax + len1 - 1;
                                            gapy++;
                                            ngapy -= siz;
        /* ignore MAXGAP when doing endgaps */
45
                                            siz = (-siz < MAXGAP | | endgaps)? -siz : MAXGAP;
                                            i1++;
                                   else if (siz > 0) { /* gap in first seq */
                                            pp[0].n[i0] = siz;
 50
                                            pp[0].x[i0] = xx;
                                            gapx++;
                                            ngapx += siz;
        /* ignore MAXGAP when doing endgaps */
                                            siz = (siz < MAXGAP | | endgaps)? siz : MAXGAP;
 55
                                            i0 + +:
                                   }
                          }
                          else
                                   break;
 60
                 }
                 /* reverse the order of jmps
                 for (j = 0, i0--; j < i0; j++, i0--)
 65
                          i = pp[0].n[i]; pp[0].n[i] = pp[0].n[i0]; pp[0].n[i0] = i;
                                                              -112-
```

```
i = pp[0].x[j]; pp[0].x[j] = pp[0].x[i0]; pp[0].x[i0] = i;
                   for (j = 0, i1--; j < i1; j++, i1--)
                             i = pp[1].n[j]; pp[1].n[j] = pp[1].n[i1]; pp[1].n[i1] = i;

i = pp[1].x[j]; pp[1].x[j] = pp[1].x[i1]; pp[1].x[i1] = i;
  5
                   if (fd > = 0)
                             (void) close(fd);
                   if (fj) {
10
                             (void) unlink(jname);
                              f_1 = 0;
                             offset = 0;
                                                                   Table 30
15
          * write a filled jmp struct offset of the prev one (if any): nw()
                                                                                                                           writejmps
         writejmps(ix)
                   int
                              ix:
20
                   char
                              *mktemp();
                    if (!fj) {
                              if (mktemp(jname) < 0) {
25
                                        fprintf(stderr, "%s: can't mktemp() %s\n", prog, jname);
                              if ((fj = fopen(jname, "w")) == 0) {
                                        fprintf(stderr, "%s: can't write %s\n", prog, jname);
30
                                        exit(1);
                    (void) fwrite((char *)&dx[ix].jp, sizeof(struct jmp), 1, fj);
                    (void) fwrite((char *)&dx[ix].offset, sizeof(dx[ix].offset), 1, fj);
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What is claimed is:

An isolated DNA molecule selected from the group consisting of: (1) a DNA molecule encoding an hIL-1Ra1 polypeptide comprising the amino acid sequence of amino acid residues from about 37 to about 203 of Figure 2 (SEQ ID NO:5); (2) a DNA molecule encoding an hlL-1Ra1 polypeptide comprising the amino acid sequence of amino acid residues from about 15 to about 193 of Figure 3 (SEQ ID NO:7); (3) a DNA molecule encoding an hIL-1Ra2 polypeptide comprising the amino acid sequence of amino acid residues from about 1 to about 134 of Figure 5 (SEQ ID NO:10); (4) a DNA molecule encoding an hIL-1Ra3 polypeptide comprising the amino acid sequence of amino acid residues from about 95 to about 134 of Figure 7 (SEQ ID NO:13); (5) a DNA molecule encoding a mIL-1Ra3 polypeptide comprising the amino acid sequence of amino acid residues from about 95 to about 134 of Figure 9 (SEQ ID NO:16); (6) a DNA molecule encoding an hIL-1Ra1L polypeptide comprising the amino acid sequence of amino acid residues from about 26 to about 207 of Figure 15 (SEQ ID NO:19); (7) a DNA molecule encoding an hIL-1Ra1S polypeptide comprising the amino acid sequence of amino acid residues from about 26 to about 167 of Figure 16 (SEQ ID NO:21); (8) a DNA molecule encoding an hIL-1Ra1V polypeptide comprising the amino acid sequence of amino acid residues from about 46 to about 218 of Figure 19 (SEQ ID NO:25); and (9) the complement of any of the DNA molecules of (1)-(8).

2. The isolated DNA molecule of claim 1 selected from the group consisting of: (1) a DNA molecule encoding an hIL-1Ra1 polypeptide comprising the amino acid sequence of amino acid residues from about 37 to about 203 of Figure 2 (SEQ ID N0:5); (2) a DNA molecule encoding an hIL-1Ra1 polypeptide comprising the amino acid sequence of amino acid residues from about 15 to about 193 of Figure 3 (SEQ ID NO:7); (3) a DNA molecule encoding an hIL-1Ra2 polypeptide comprising the amino acid sequence of amino acid residues from about 1 to about 134 of Figure 5 (SEQ ID NO:10); (4) a DNA molecule encoding an hIL-1Ra3 polypeptide comprising the amino acid sequence of amino acid residues from about 95 to about 134 of Figure 7 (SEQ ID NO:13); (5) a DNA molecule encoding a mIL-1Ra3 polypeptide comprising the amino acid sequence of amino acid residues from about 95 to about 134 of Figure 9 (SEQ ID NO:16); and (6) the complement of any of the DNA molecules of (1)-(5).

3. The isolated DNA molecule of Claim 1 selected from the group consisting of: (1) a DNA molecule encoding an hIL-1Ra1L polypeptide comprising the amino acid sequence of amino acid residues from about 26 to about 207 of Figure 15 (SEQ ID NO:19); (2) a DNA molecule encoding an hIL-1Ra1S polypeptide comprising the amino acid sequence of amino acid residues from about 26 to about 167 of Figure 16 (SEQ ID NO:21); (3) a DNA molecule encoding an hIL-1Ra1V polypeptide comprising the amino acid sequence of amino acid residues from about 46 to about 218 of Figure 19 (SEQ ID NO:25); and (4) the complement of any of the DNA molecules of (1)-(3).

4. The isolated DNA molecule of Claim 3 selected from the group consisting of: (1) a DNA molecule encoding an hIL-1Ra1L polypeptide comprising the amino acid sequence of amino acid residues from about 1 to about 207 of Figure 15 (SEQ ID NO:19); (2) a DNA molecule encoding an hIL-1Ra1S polypeptide comprising the amino acid sequence of amino acid residues from about 1 to about 167 of Figure 16 (SEQ ID NO:21); (3) a DNA molecule encoding an hIL-

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1Ra1V polypeptide comprising the amino acid sequence of amino acid residues from about 1 to about 218 of Figure 19 (SEQ ID NO:25); and (4) the complement of any of the DNA molecules of (1)-(2).

- The isolated DNA molecule of Claim 2 selected from the group consisting of: (1) a DNA molecule encoding an hIL-1Ra3 polypeptide comprising the amino acid sequence of amino acid residues from about 34 to about 155 of Figure 7 (SEQ ID NO:13); (2) a DNA molecule encoding a mIL-1Ra3 polypeptide comprising the amino acid sequence of amino acid residues from about 34 to about 155 of Figure 9 (SEQ ID NO:16); and (3) the complement of any of the DNA molecules of (1)-(2).
- 6. The isolated DNA molecule of Claim 5 selected from the group consisting of: (1) a DNA molecule encoding an hIL-1Ra3 polypeptide comprising the amino acid sequence of amino acid residues from about 2 to about 155 of Figure 7 (SEQ ID NO:13); (2) a DNA molecule encoding a mIL-1Ra3 polypeptide comprising the amino acid sequence of amino acid residues from about 2 to about 155 of Figure 9 (SEQ ID NO:16); and (3) the complement of any of the DNA molecules of (1)-(2).
- 7. The isolated DNA molecule of Claim 1 selected from the group consisting of: (1) a DNA molecule which encodes an hIL-1Ra1 polypeptide, and which comprises the nucleic acid sequence of nucleotide positions from about 118 to about 618 in the sense strand of Figure 2 (SEQ ID NO:4); (2) a DNA molecule which encodes an hIL-1Ra1 polypeptide, and which comprises the nucleic acid sequence of nucleotide positions from about 145 to about 681 in the sense strand of Figure 3 (SEQ ID NO:6); (3) a DNA molecule which encodes an hIL-1Ra2 polypeptide, and which comprises the nucleic acid sequence of nucleotide positions from about 96 to about 497 in the sense strand of Figure 5 (SEQ ID NO:9); (4) a DNA molecule which encodes an hIL-1Ra3 polypeptide, and which comprises the nucleic acid sequence of nucleotide positions from about 283 to about 402 in the sense strand of Figure 7 (SEQ ID NO:12); (5) a DNA molecule which encodes a mIL-1Ra3 polypeptide, and which comprises the nucleic acid sequence of nucleotide positions from about 427 to about 546 in the sense strand of Figure 9 (SEQ ID NO:15); (6) a DNA molecule which encodes an hIL-1Ra1L polypeptide, and which comprises the nucleic acid sequence of nucleotide positions from about 79 to about 624 in the sense strand of Figure 15 (SEQ ID NO:18); (7) a DNA molecule which encodes an hIL-1Ra1S polypeptide, and which comprises the nucleic acid sequence of nucleotide positions from about 79 to about 504 in the sense strand of Figure 16 (SEQ ID NO:20); (8) a DNA molecule which encodes an hIL-1Ra1V polypeptide, and which comprises the nucleic acid sequence of nucleotide positions from about 208 to about 726 in the sense strand of Figure 19 (SEO ID NO:24); and (9) the complement of any of the DNA molecules of (1)-(8).
- The isolated DNA molecule of Claim 7 selected from the group consisting of: (1) a DNA molecule which encodes an hIL-1Ra1 polypeptide, and which comprises the nucleic acid sequence of nucleotide positions from about 118 to about 618 in the sense strand of Figure 2 (SEQ ID NO:4); (2) a DNA molecule which encodes an hIL-1Ra1 polypeptide, and which comprises the nucleic acid sequence of nucleotide positions from about 145 to about 681 in the sense strand of Figure 3 (SEQ ID NO:6); (3) a DNA molecule which encodes an hIL-1Ra2

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polypeptide, and which comprises the nucleic acid sequence of nucleotide positions from about 96 to about 497 in the sense strand of Figure 5 (SEQ ID NO:9); (4) a DNA molecule which encodes an hIL-1Ra3 polypeptide, and which comprises the nucleic acid sequence of nucleotide positions from about 283 to about 402 in the sense strand of Figure 7 (SEQ ID NO:12); (5) a DNA molecule which encodes a mIL-1Ra3 polypeptide, and which comprises the nucleic acid sequence of nucleotide positions from about 427 to about 546 in the sense strand of Figure 9 (SEQ ID NO:15); and (6) the complement of any of the DNA molecules of (1)-(5).

- 9. The isolated DNA molecule of Claim 7 selected from the group consisting of: (1) a DNA molecule which encodes an hIL-1Ra1L polypeptide, and which comprises the nucleic acid sequence of nucleotide positions from about 79 to about 624 in the sense strand of Figure 15 (SEQ ID NO:18); (2) a DNA molecule which encodes an hIL-1Ra1S polypeptide, and which comprises the nucleic acid sequence of nucleotide positions from about 79 to about 504 in the sense strand of Figure 16 (SEQ ID NO:20); (3) a DNA molecule which encodes an hIL-1Ra1V polypeptide, and which comprises the nucleic acid sequence of nucleotide positions from about 208 to about 726 in the sense strand of Figure 19 (SEQ ID NO:24); and (4) the complement of any of the DNA molecules of (1)-(3).
- 10. The isolated DNA molecule of Claim 9 selected from the group consisting of: (1) a DNA molecule which encodes an hIL-1Ra1L polypeptide, and which comprises the nucleic acid sequence of nucleotide positions from about 4 to about 624 in the sense strand of Figure 15 (SEQ ID NO:18); (2) a DNA molecule which encodes an hIL-1Ra1S polypeptide, and which comprises the nucleic acid sequence of nucleotide positions from about 4 to about 504 in the sense strand of Figure 16 (SEQ ID NO:20); (3) a DNA molecule which encodes an hIL-1Ra1V polypeptide, and which comprises the nucleic acid sequence of nucleotide positions from about 73 to about 726 in the sense strand of Figure 19 (SEQ ID NO:24); and (4) the complement of any of the DNA molecules of (1)-(3).
- 11. The isolated nucleic acid molecule of Claim 8 selected from the group consisting of: (1) a DNA molecule comprising the nucleic acid sequence of nucleotide positions from about 103 to about 681 in the sense strand of Figure 3 (SEQ ID NO:6); (2) a DNA molecule comprising the nucleic acid sequence of nucleotide positions from about 100 to about 465 in the sense strand of Figure 7 (SEQ ID NO:12); (3) a DNA molecule comprising the nucleic acid sequence of nucleotide positions from about 244 to about 609 in the sense strand of Figure 9 (SEQ ID NO:15); and (4) the complement of any of the DNA molecules of (1)-(3).
- 12. The isolated nucleic acid molecule of Claim 8 comprising (a) the complete DNA sequence in the sense strand of Figure 2 (SEQ ID NO:4), Figure 3 (SEQ ID NO:6), Figure 5 (SEQ ID NO:12), or Figure 9 (SEQ ID NO:15), or (b) the complement of (a).
- 13. An isolated nucleic acid molecule encoding an 1L-1lp polypeptide, comprising DNA hybridizing to the complement of a nucleic acid sequence selected from the group consisting of: (1) the nucleic acid sequence consisting of nucleotide positions from about 238 to about 465 in the sense strand of Figure 7 (SEQ ID NO:12); (2) the nucleic acid sequence consisting of nucleotide positions from about 427 to about 609 in the sense strand of Figure 9 (SEQ ID

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NO:15); and (3) the nucleic acid sequence consisting of nucleotide positions from about 114 to about 135 in the sense strand of Figure 15 (SEQ ID NO:18).

- An isolated nucleic acid molecule comprising (a) a DNA molecule encoding a polypeptide selected from the group consisting of: (1) a polypeptide comprising the entire amino acid sequence encoded by the longest open reading frame in the cDNA insert in the vector deposited as ATCC Deposit No. 203588; (2) a polypeptide comprising the entire amino acid sequence, or the entire amino acid sequence excluding the 36 N-terminal amino acid residues of such sequence, encoded by the longest open reading frame in the cDNA insert in the vector deposited as ATCC Deposit No. 203587; (3) a polypeptide comprising the entire amino acid sequence encoded by the longest open reading frame in the cDNA insert in the vector deposited as ATCC Deposit No. 203586; (4) a polypeptide comprising the entire amino acid sequence, or the entire amino acid sequence excluding the N-terminal amino acid residue of such sequence, encoded by the longest open reading frame in the cDNA insert in the vector deposited as ATCC Deposit No. 203589; (5) a polypeptide comprising the entire amino acid sequence, or the entire amino acid sequence excluding the N-terminal amino acid residue of such sequence, encoded by the cDNA insert in the vector deposited as ATCC Deposit No. 203590; (6) a polypeptide comprising the entire amino acid sequence, or the entire amino acid sequence excluding the 34 N-terminal amino acid residues of such sequence, encoded by the longest open reading frame in the cDNA insert in the vector deposited as ATCC Deposit No. 203846; (7) a polypeptide comprising the entire amino acid sequence encoded by the longest open reading frame in the cDNA insert in the vector deposited as ATCC Deposit No. 203855; and (8) a polypeptide comprising the entire amino acid sequence, or the entire amino acid sequence excluding the 45 N-terminal amino acid residues of such sequence, encoded by the longest open reading frame in the cDNA insert in the vector deposited as ATCC Deposit No. 203973; or (b) the complement of the DNA molecule of (a).
- An isolated nucleic acid molecule comprising (a) DNA encoding the IL-1lp polypeptide of Claim 21, or (b) the complement of the DNA of (a).
  - A vector comprising the nucleic acid of Claim 1. 16.
- The vector of Claim 16 operably linked to control sequences recognized by a host 17. cell transfected with the vector.
  - A host cell comprising the vector of Claim 16.
- A process for producing an IL-11p polypeptide comprising the steps of: (1) culturing a host cell comprising the DNA molecule of claim 15 under conditions suitable for expression of the IL-11p polypeptide encoded by the DNA molecule; and (2) recovering said IL-11p polypeptide from the cell culture.
  - An isolated IL-11p polypeptide encoded by the nucleic acid molecule of Claim 1. 20.
- An isolated IL-11p polypeptide selected from the group consisting of: (1) an hIL-1Ra1V polypeptide consisting of an amino acid sequence having at least an 80% sequence identity to the sequence of amino acid residues from about 46 to about 218 of Figure 19 (SEQ ID NO:25); (2) an hIL-1Ra3 polypeptide consisting of an amino acid sequence having at least an 80% sequence identity to the sequence of amino acid residues from about 95 to about 134 of

Figure 7 (SEQ ID NO:13); and (3) a mIL-1Ra3 polypeptide consisting of an amino acid sequence having at least an 80% sequence identity to the sequence of amino acid residues from about 95 to about 134 of Figure 9 (SEQ ID NO:16).

An isolated IL-11p polypeptide selected from the group consisting of: (1) an hIL-22. 1Ra1 polypeptide comprising amino acid residues from about 37 to about 203 of Figure 2 (SEQ ID NO:5); (2) an hIL-1Ra1 polypeptide comprising amino acid residues from about 15 to about 193 of Figure 3 (SEQ ID NO:7); (3) an hIL-1Ra2 polypeptide comprising amino acid residues from about 1 to about 134 of Figure 5 (SEQ ID NO:10); (4) an hIL-1Ra3 polypeptide comprising amino acid residues from about 95 to about 134 of Figure 7 (SEQ ID NO:13); (5) a mIL-1Ra3 polypeptide comprising amino acid residues from about 95 to about 134 of Figure 9 (SEQ ID NO:16); (6) an hIL-1RalL polypeptide comprising amino acid residues from about 26 to about 207 of Figure 15 (SEQ ID NO:19); (7) an hIL-1Ra1S polypeptide comprising amino acid residues from about 26 to about 167 of Figure 16 (SEQ ID NO:21); and (8) an hIL-1Ra1V polypeptide comprising amino acid residues from about 46 to about 218 of Figure 19 (SEQ ID NO:25).

- The isolated IL-1lp polypeptide of Claim 22 selected from the group consisting of: (1) an hIL-1Ra1L polypeptide comprising amino acid residues from about 1 to about 207 of Figure 15 (SEQ ID NO:19); (2) an hIL-1Ra1S polypeptide comprising amino acid residues from about 1 to about 167 of Figure 16 (SEQ ID NO:21); and (3) an hIL-1Ra1V polypeptide comprising amino acid residues from about 1 to about 218 of Figure 19 (SEQ ID NO:25).
- The isolated polypeptide of Claim 22 selected from the group consisting of: (1) an hIL-1Ra3 polypeptide comprising amino acid residues from about 34 to about 155 of Figure 7 (SEQ ID NO:13); and (2) a mIL-1Ra3 polypeptide comprising amino acid residues from about 34 to about 155 of Figure 9 (SEQ ID NO:16).
  - The polypeptide encoded by the DNA molecule of (a) in claim 14. 25.
- The IL-1lp polypeptide of Claim 22 that comprises a native amino acid sequence 26. of the IL-11p fused at its C-terminus or N-terminus to a heterologous amino acid sequence.
- 27. The IL-11p polypeptide of Claim 26, wherein said heterologous amino acid sequence is an epitope tag sequence.
- The IL-11p polypeptide of Claim 26, wherein said heterologous amino acid sequence is a Fc region of an immunoglobulin.
  - An antibody which specifically binds to the IL-11p polypeptide of claim 22. 29.
  - The antibody of Claim 29, wherein said antibody is a monoclonal antibody. 30.

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- GODICORGOS ANGOCTICCA, GOTINICOTO ACCALCUTO ANAGICIGA AGCINCIDOC CTACAGAMAS TINCTAGIOS CCIMAAGCIG CCOTOCICCO TICOGRÁGOS COANTAGEAC ISCOTOGRAC TITCAGACIC TGANTAKOS GAIGHCTITIC ANTGAICACS GANTIFOGAC start insert
  - possible splice donor IOI TRANGITACT GCIGCTGITG GAGTACAACT ICCCTATMCA AAACAACTGC CAGCACCTA AGACACTCA CACCTGCAGA <u>GIGGGC</u>TTGA GAAGATTTG ACTACAATGA CGACGARAAC CTCATGFTGA AGGGARATCT ITTGITGACG GICGIGGGAAT ICTGGTGAG GIGGAAGTCT CACCGGAACT CITTCTAAAC F E × H 0 ACTACAATGA CGACGACAAC CTCATGTTGA AGGGATATCT TTTGTTGACG U N N ы Δ, ы 1 MLL
- AGINCICGCT CTYGIGGIGN ATTCTCCTAT CACTIGNICA GACGINCACT CTGCGACTCT AGGATACAGT CCGACACTAT CCTCCCTTIG 201 GGGTCAAGGA TCATGAGGGA GAAGACCACT TAAGAGGATA GTGAACTAGT CTGCATGTGA GACGCTGAGA TCCTATGTCA GGCTGTGATA GGAGGGAAAC CCCAGTTCCT
  - 301 AGAMACCARA GGARAGACA GCTITAAGAA GGGCTTAAGA GCCACCCACC CATTCTTGAC AGTCACTGGC CCAGCCTGGG GGCCCCTGTT CTTTATCAAA TCTTTGGTT CCTTTCTTGT CGAAATTCTT CGCGAATTCT CGGTGGGTGG GTAAGAACTG TCAGTGACCG GGTCGGACCC CCGGGGACAA GAAATACTTT
    - 401 CANGIGCTG AGTOTITGC RANGITCAA AGGIGAAGAA CITAAACCCG AAGAAAITGA GCATICAIGA CCAGGATCAC AAAGTACTGG TOCTGGACTC GTCAACGAAC TGGGGAAGG TGTGCAGGIT ICCACTTCTT GAATTIGGGC TTCTITAAGT CGTAAGTACT GGTGCTAGTG TTTCATGACC AGGACCTGAG O D LNPKKFSIHD ~85066.AH1282.Asc.f, 5'Tag: TTTCCCTTTGGCGCGCC
      - '85066.AH1284.NSi.f., 5'Tag: TTTCCCTTTATGCATCAGATGACGATGACGAAAA 'possible splice acceptor 'ORF
- SOI TGGGAATCTC ATAGCAGTIC CAGAFAAAAA CTACATACGC CCAGAGATCT TCTTTGCATT AGCCTCATCC TTGAGGTCAG CCTCTGGGGA ACCCTINGAG TATCGTCAAG GTCTATTTTT GATGTATGCG GGTCTCTAGA AGAAACGTAA TCGGAGTAGG AACTCGAGTC GGAGACGCCT S a FAL P E F Y I N ρ, I A V
- 601 CCGATTCTCC TGGGGGTCTC TARAGGGGAG TTTTGTCTCT ACTGTGACAA GGATAAAGGA CAAAGTCATC CATCCCTTCA GCTGAAGAAG GAGAAACTGA L K K GOCTAAGAGG ACCCCCAAGA ATTCCCCTC AAAACAGAGA TGACAKTTTT CCTATTTCCT GITTCAGTAG GTAGGGAAGT P 1 L L G V S K G E F C L Y C D K D K G Q S H P S L Q P I L L G V S K G E F C L Y C D K D K G Q S H P S L Q 57

901 ААЛЕСТВАВ ТЕЛЕССЕЛЬ ТВАБОТОЛО ПОВОЗТИВОВЛА СТОСССЕЛТ БАЛСССТИТ СТОСТВАТИТ ТВАРСТВАТИ ОТВИТИТИТО ТВОТИТОВЛЕ ТИТОСЛЕТЬ АСТОСАВТЕ СТАЛІТИТИ ОБОТИТОВАС СТОСОВОТО В V S D V S R A B H  $\tilde{S}$  P S E V S D V701 TGANGCTGGC TGCCCAAAAG GAATCAGCAC GCCGGCCCTT CATCTTTAT AGGGTCAGG TGGGCTCCTG GAACATGCTG GAGTCGGCGG CTCACCCGG ACTICOACCE ACEGETITIC CTTAGECTE GESCESSAA STAGAAAATA TCCCAAGTC ACECSAGAC CTTGTAGAC CTCAGCGGC GAGTGGGGCC BOI ARGETICATO TOCACOTOCY GCARITORA. TORGOCYCHY GOGGOGGICA ATACHTORA GARCAGGIAA CACATGGAAT TYTCATTCA ACCAGTYTOC TACCAAGTAG AGOTGGAGGA COTTAACAT ACTGGGACA CCCCACTOC TATTINAACT CYTGTOCYTY GYGTAACTTA AAAGTAAAGT TOGTCAAAGG HIEF N R N RAOV GVTDKFE I F 74 14 ЕР ESAR N C N CTSC A Q K K L A 124 W F I 91

^85066.AH1283.r ^85066.AH1285.Not.r, 5'Tag: TTTCCTTTGCGGCCGCTTA

1001 CTCACT GAGTGA

			S
JC	A.G	a	nsert
AGC	TCG	A	
PTC	PAG.	Ø	
AA	C TTAAGTCGA	z	
50500	30000	ď	
999	SSS	æ	
73	ACG	æ	
CCI	CGA	ы	
CAA	GTT	×	
ACGA	TGCT	Ω	
E E	AC.	П	
CGA	SCI	Ω	
AGA	TCT.	Ω	
Ā	Ħ	×	
CTAC	GATG	×	^flag
췵	ACT	Ω	4
TGC	ACG	×	
AGT	BAACAAC CTCGACGTCA ACGACTGATG	>	
TGC	3AC	4	
GAGC	CICC	¥	
FIG.	AAC	O	
PTG	NO.	>	
DI	3AG	ч	
9	8	4	
CL	3GA	ü	
BAT	CTA	н	
TCT	GA AGACTAGGAT	J	
ACT	IGA	ы	
TGC	ACG	ď	
IGIC	ACAG	S	4
TAATTCACCA	ATTAAGTGGT ;	М	400
-		H	

101 TTTGCAGAGG TCCAAAGGTG AAGAACTTAA ACCCGAAGAA ATTCAGCATT CATGACCAGG ATCACAAAGT ACTGGTCCTG GACTCTGGGA ATCTCATAGC TGACCAGGAC CTGAGACCCT TAGAGTATCG T V L TOGGCTTCTT TAAGTCGTAA GTACTGGTCC TAGTGTTTCA н к 0 Ωн F S I M M TTCTTGAATT KNLN AAACGTCTCC AGGTTTCCAC РК

CITCAGGCIA AGAGGACCCC GCGGAGAAAG GAAGTCCGAT TCTCCTGGGG CGCCTCTTTC A E K 201 AGTICCAGAI AAAAACTACA TACGCCCAGA GAICTICITI GCAITAGCCI CAICCITGAG CICAGCCICI TCAAGGICTA ITITIGAIGI AIGCGGGICI CTAGAAGAAA CGIAATCGGA GTAGGAACTC SAGTCGGAGA SAS S ALAS E E ELI Cu æ

LAAQ CHGTHCCTAT ITCCHGTTTC AGTAGGTAGG GAAGTCGACT TCTTCCTCTT TGACTACTTC GACCGACGGG 301 GTCTCTAAAG GGGAGTTTTG TCTCTACTGT GACAAGGATA AAGGACAAAG TCATCCATCC CTTCAGCTGA AGAAGGAGAA ACTGATGAAG E E × LOLK H P S 3 0 9 DKDK CAGAGATTTC CCCTCAAAAC AGAGATGACA U 7 E P C

101 ARARGRATE AGENEGEGG COUTCATET TTRATAGGG TEAGGTGGG TECTGGRAEA TGCTGGAGTE GGGGGTEAE CCGGARGGT TEATETGEAE CCGCCGAGTG GGGCCTACCA AGTAGACGTG N C M A A H TITICCTIAG ICGIGCGGCC GGGAAGIAGA AAAIAICCCG AGICCACCG AGGACCIIGI ACGACCICAG SWNM o ∧ ở A R R

CCTTTGTGTA ACTTAAAAGT AAAGTTGGTC AAACGTTTGG ACTTTACTCG TITCAACCAG TITGCAAAGC TGAAATGAGC 501 CTCCTGCAAT TGTAATGAGC CTGTTGGGGT GACAGATAAA TTTGAGAACA GGAAACACAT TGAATTTTCA GAGGACGITA ACAITACTCG GACAACCCCA CIGICTATIT AAACICTIGI F E N R 5 CNE 165

601 CCCAGTGAGG TCAGCGATTA GGGTACCAGT CGAGTCTAGA GGATCCCGGG GGGTCATCC AGTGGTAAA CCCATGGTCA GTGAGATCT CTAGGGCCC 198 F S E V S D O

^inserts ends

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- GCGCTGGCAC CCTAAAGCTG GGATTTCGAC TTACTAGTGC GGTTATCGTG ACGCACCTTG AAAGTCTGAG AGCTACTGCC CTACAGAAAG CCAATAGCAC TGCGTGGAAC TTTCAGACTC TCGATGACGG GATGTCTTTC GGCACGAGGC AAGCCTTCCA
- GTGAAGACT TAAACCCGAA z CACTTCTTGA N K CACCTTCAGA GTGGAAGTCT £L, <u>-</u> TCCCTATAGA AAACAACTGC CAGCACCTTA AGACCACTCA GTCGTGGAAT TCTGGTGAGT Ħ e e L K ; 0 TTTGTTGACG O z 2 CTCATGTTGA AGGGATATCT Н GAGTACAACT Y N 101 TGATGTTACT GCTGCTGTTG CGACGACAAC 1 ü ACTACAATGA T E
  - TCTCTAGAAG AGAGATCTTC GATAAAACT ACATACGCCC TGTATGCGGG CTATTTTGA AGGATCACAA AGTACTGGTC CWGGACTCTG GGAATCTCAT AGCAGTTCCA TCCTAGAGTT TCATGACCAG GACCTGAGAC CCTTAGAGTA TCGTCAAGGT TAAGTACTGG ATTCATGACC GAAATTCAGC CTTTAAGTCG 201
    - М AACAGAGATG ACACTGTTCC TTGTCTCTAC I R P CLY D K N Y GALTCTCCTG GGGGTCTCTA AAGGGGAGTT CCCCAGAGAT TTCCCCTCAA ш AV S A 9 CTAAGAGGAC 1 ILL 2 TCTGCGGAGA AAGGAAGTCC AGACGCCTCT TTCCTTCAGG Ŋ Ω O SAEK Þ \ \ GAGCTCAGCC CTCGAGTCGG × S Ξ Ω ξŊ 301 TTTGCATTAG CCTCATCCTT GGAGTAGGAA S H AAACGTAATC 29 34
- GCCGGGAAGT AGAAAATATC CGGCCCTTCA GGGTTTTCCT TAGTCGTGCG ATCAGCACGC ac) 4 o CCCAAAAGGA 0 AAGCTGGCTG TTCGACCGAC æ K GAAACTGATG CTTTGACTAC I. × ACTICITICET . K K E TGAAGAAGGA TCCCTTCAGC AGGGAAGTCG 0 o 401 ATAAAGGACA AAGTCATCCA TTCAGTAGGT V. TATTTCCTGT 101
- CCACTGTCTA н Þ CACCCCGGAI GSITCATCIG CACCICCIGC AAIIGIAAIG AGCCTGIIGG GIGGGGGAC THAACAITAC ICGGACAACC > TTAACATTAC υ 2 GTGGAGGACG ı s CCAAGTAGAC GIGGGGCCIA M S G. GTCGGCGGCT CAGCCGCCGA SAA ACATGCTGGA TGTACGACCT × GGCTCAGGTG GGCTCCTGGA CCGAGTCCAC CCGAGGACCT 3 S 501 134
- TCGGGGTCAC TCCAGTCGCT AATCCTTTGA CGGGGTAACT S P S E V S D 0 AGCCCCAGTG AGGTCAGCGA AGCTGAAATG TCGACTTYAC × ы æ CAGTTTGCAA GTAACTTAAA AGTAAAGTTG GTCAAAGGTT > TCATTTCAAC ø CATTGAATTT AAATTTGAGA ACAGGAAACA TTTABACTCT TGTCCTTTGT × 601 167

TTAGGAAACT

ACGCCTTCCT CGCTAATTTG AACTAATTGT ATAAAAACAC CAAACCTGCT TGCGGAAGGA GCGATTAAAAC TTGATTAAACA TATTTTTGTG GTTTGGACGA 701

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1 COAGOCCOA GCNICCCCAC CATGAATIT GITCACACAA GTCGAAAGGT GAAGAGCTTA AACCCGAAGA AATCAGCAT TCATGACCAG GATCACAAAG GSTCCGSGTT GSNAGGGSTG GTACTTAAAA CAAGTGTGT CAGCTTTCCA CTTCTCGAAT TTGGGCTTCT TTAAGTGSTA AGTACTGSTC CTAGTGTTTC

101 IACTOGOCTO GACTOTOGRA MOTOCRARAO AGTICCARAR AAAAACTACA TACOCCOLGA GATOTICTIT GCATUAGOCT CATOCTIGAG CTCAGOCTOT ATGACOGGAC CTGAGACCCT TAGAGTATCG TCAAGGTOTA TTTTTGATOT ATGAGGGTOT CTAGAAGAAA CGTAATGGGA GTAGGAACTO GAGTGGAGA

201 GCGGAGAAAG GAAGTCCGA<sub>R</sub> TCTCCTGGGGG GTCTCTAAAG GGGAGTTTTG T<u>CTCTACTGT GACAAGGARA AAGGACAAAG TCATCCATCC CTTCAGCTGA</u> COCCUCTIVE CITCHOSCIA AGRIGACIACCE CAGAGATUTE CECTCAAAAC AGAGATGAACA CIRSTICETAA TVECTORITYE AGIAAGIAGO GAAGTGAACT I L L G V S K G B F C L Y C D K D K G Q S H P S L Q L K ^84664.pl

301 AGAAGGAGAA ACTGATGAAG CTGACTGCCC AAAAGBAATC AGCACGCGG CCCTTCATCT TTTATAGGGC TCAGGTGGGC TCCTGGAACA TGCTGGAGC PETICOTOTI TEACTACITO GACCGACGES ITTICCITAS ICGIGOGGCC GESAAGIAGA AAAINICCOG AGICCACCCG AGGACCITGI ACGACCITAG S W N M 0 0 X R FF A R R M LAAQ 29

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401 GGGGGCTCAC CCCGGATGGT TCATCTGCAC CTCCTGCAAT TGTAATGAGC CTGTTGGGGT GACAGATAAA TTTGAGGAAGA GGAAACACAT TGAATTTCA COGCOGAGTG GGGCCTACCA AGTAGACGTG GAGGACGTTA ACATTACTCG GACAACCCCA CTGTCTATTT AAACTCTTGT CCTTTGTGTA ACTTAAAAGT T D K A G A CNEP SCN T C I E W E 62 501 THICAACCAG THTGCAAAGC TGAAATGAGC CCCAGTGAGG TCAGCGAFTTA GGBAAGGGC CCATTGAAGG CCTTCCTCGC TAATTTGAAC TAATTGTATA AAAGTTGGTC CAATGATTGTAG GGAAGGAGGG ATTAAACTTG ATTAACATAT ~84664.rl g S PSEV N N ک ط 95

601 AAAACCCCAA ACCTGCTCAC TAAAAAAA TTTTGGGGTT TGGACGAGTG ATTTTTTT 9/869566 PCT/US99/30720

~92929.AHJ CAGIATGITG TGACGGAAAC TIAATTATA TTTACTGACT S Y N T A F E L N I N D O 1 OFCARCERE GESTECGAAG CROSTICAAC CACANTRAG FECCETGAG TGTAGATAAA GACCETTET TGCGAGGTGE TGAGAGAACE ACACT<u>AGAG</u> CAGTIGGTG GGCAGGTTC GACGACCTG GTGCTAAGTC AGGGGACCTG ACATCTATTT CTGGGAAAGA ACGGTCCAGG ACTCTGTTGG TGTGATACTC ACTCGTTCCG GGAAGGGCCT GTCATACAAC ACTGCCTTTG AATTAAATAT AAATGACTGA 501 ACTOROCCIA GAGINGGENG CITGGECTIT GECTRANGT TECTOSTEC CANGEGITT TOGECIACAT TITCTRAGG TANTITICAG GCTOSTGCTG FGAGIOGSAT CYCCACCGTC GAACCAGAAA CAGAATITCA AAGACCAAGG GTRACACAA AGAGATGTA AAAGATCAC GATAAAAGTG GGACCACGAC AGACAGGAGC AAGGCTGCTG TTATCATCTC ATTITATAAT GAAGAAGAAG CAATTACTTC ATAGCAACTG AAGAACAGGA TGTGGCCTCA GAAGCAGGAG TORGECTICS TICCARGAR AATAGRAGAS TARAATATTA CITCITCITC SITAAIGAAS TARCSITGAC TICITGICCT ACACCGGAGI CITCSICCTC 101 AGCTGGGTGG TATAAGGCTG TCCTCTCAAG CTGGTGCTGT GTAGGCCACA AGGCATCTGC ATGAGTGACT TTAAGACTCA AAGACCAAAC ACTGAGCTTT TOGROCCACC AFAITOCGAC AGGAGAGITO GACCACGACA CATOCGGIGI TOCGIAGACG TACTCACTGA AATTCTGAGI TTCTGGITIG TGACTCGAAA 92929. AH1421. Asc.f, 5'Tag: AAAGGGAAAAGGCGCGCC^ AGGCACTCCA GOAGACGCTG ATGGTGGAGG AAGGGCCGTC TATCAATCAA TCACTGTTGC TGTTATCACA TGCAAGTATC CAGAGGCTCT ACGITCATAG GICTCCGAGA INGIGAGAAG GINGGAGAAC AGCCCACAIT GCAGCIAAAA TCGGGTGTAA CGTCGATTTT GAGGTGGGAA CTCAGACACC E A L α CKY CTCCACCCTT ST TCCGTGAGGT CCTCTGCGAC TACCACCTCC TTCCCGGCAG ATAGTTAGTT AGTGACAACG ACAATAGTGT TRAGGICTIT ACACAACAT AACACITTC CAACCICITG CTGGTAGGAC GACCATCCTG E 65 ΙI E C CGTGCCAAGA GCACGGTTCT RAKT GCCTCCTCCA AGAGAGACCA GCCCATCATT CTGACTTCAG AACTTGGGAA CGGGTAGTAA GACTGAAGTC TTGAACCCTT T C M M ... Y Q S I CCATTTATTT GGGAATCCAG AATCCAGAAA TGTGTTTGTA CCCGAGCCCG TGAAACCCTT CCTTTTCTAC GGGCTCGGGC ACTTTGGGAA GGAAAAGATG r Z LFY co ပ NPEM D. R A V H GGTAAATAAA CCCTTAGGTC PEPV CGGAGGAGGT TCTCTCTGGT 0 1 0 Д œ CATACCGGTT GTATGGCCAA I Y L G D 201 AGAGGGGATC TCTCCCCTAG 301 TCATGGATCT AGTACCTAGA 401 CTGGTTCATT GACCAAGTAA R G D c 601 101 36

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СТСТИОВОВ, ТОВЛЕДСКИЕ СВОТСТЕТИИ ВОБИТОСТВИ ТОТОТОТИИ ВОВРОСТЕТ ТОВСЕТИВО ВОСТОТВИВ СМАМАНИИТ ОЗВОДНОСТ ВСТИТИТИ ОТСЛОВАМИ ТЯСНОВОЕТ ВОВИСЯСНИЕ ОСПОСЛЕТИ ТСТГОВАЮВ АТСОТИЛЕТС ТОВЛАСИТЕ СПЕТИТИТИ 801 CITCTAGGG TGGGTATGAA GATGCTTCAG AGCTCATGCG CGTTACCCAC GATGGCATGA CTAGCACAGA GCTGATCTCT GTTTCTGTTT TGCTTTATTC GAMGATICCIC ACCONTACTT CTACGAAGTIC TCGAGTACGC GCAATGGGTG CTACCGTACT GATCGTGTCT CGACTAGAGA CAAAGACAAA ACGAAATAAG 901

NGANCECCAE CENTRETTET ACGANGTETE GAGTACGEGE AMIGGETGET ACCGINETGA TEGTETETEG ACTAGAGAEA AAGACAAAAE GAAATAAGGG 1101 TCTTGGGAFG ATRICATCCA GTCTTTATAT GTTGCCAATA TACCTCATTG TGTGTAATAG AACCTTCTTA GCATTAAGAC CTTGTAAACA AAAATACT TCTTGGGGTG GĞTATGAAGA TGCTTCAGAG CTCATGCGGC TTACCCACGA TGGCATGACT AGCACAGAGC TGATCTCTGT TTCTGTTTTG 1001

1201 TICOGTIAAG TUAAACATI, TITGECCIDA, TICDAAUGIG TAATCTIAAA GITDAADAA, CITTGEGTAT TUAUAUATI, ATAAAGCUA, AACTGATATA AACACATIC AATTHAGDA AACAGGAT AACHTACAC ATHGAATIT CAATTHATIT GAACACATA AATATHITI TATITGATT TIGACTHIMI NGAROCCTAC TATAGTAGGT CAGARATATA CAROGGTTAT ATGGAGTARC ACACATTATC TTGGARGAAT CGTAATTCTG GAACATTTGT TTTTATTAAG

1301 AAATAAAGAA AGAGTAAACT TTTATTTCTT TCTCATTTGA

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1 AAGCIGCIGG AGCAACBATT CAGTCCCCTG GACTGIAGAI AAAGACCCTT ICTTGCCAGG IGCIGAGACA ACCACACTAI GAGAGGCACI CCAGGAAGACG TICGACGACC TCGGTGCTAA GTCAGGGGAC CTGACATCIA TITCTGGGAA AGAACGGTCC ACGACTCIGT TGGTGTGATA CTCTCCGTGA GGTCCTGTC

101 CHARTEGROS AGANGESCC STUTATEAR CARTESTRET ROTHSTEAR AGANGEAGE RECERCAGE TUTTGAGGA GGUAGAGGS AFCCCATATA GAUTACORC TUTTGAGGA CAGANGESC COTUTCOCC TAGGGGAAAA

201 TTTGGGAATC CAGAATCCAG AAATGTGTTT GTATTGTGAG AAGGTTGGA AAACCCTTAG GTCTTAGGTC TTTACACAAA CATAACACTC TTCCAACCT

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	ATGGTCCTGA TACCAGGACT M V L S	GTGGGGCGCT CACCCCGCGA G A L	1 AFGREGARIA GROGOGOGO TROCTICORA ARGAGARAT COCONTRATA RECENTATA RECENTATOR TOCATORAGE GROCATORAGE TROCTICORA COCONTRACT COCONDACA TOCATORAGE TROCTICORA COCONTRACT COCONDACA COCONTRACT COCONDACA TOCATORAGE ACCONTRACTOR ACCONTRACT A GOOD LE A GOOD	ATGAAGGACT TACTTCCTGA M K D S	CGGCATTGAA GCCGTAACTT A L K	GGTGCTTTAT CCACGAAATA V L Y	CTGCATAATA GACGTATTAT L H N N	ACCAGCTTCT TGGTCGAAGA Q L L	AGCTGGAGGG TCGACCTCCC A G G	CTGCATGCAG GACGTACGTC L H A G	
35	GGAAGGTCAT CCTTCCAGTA K V I	TAAAGGTGAA ATTTCCACTT K G E	101 GGANGGTCAN TANAGGTGAA GAGATCAGCG TGGTCCCCAA TCGGTGGCTG GANGCCAGCC TGTCCCCGGT CATCCTGGGT GTCCAGGGGG GAAGCCAGTG CTTCCAGTT CTCTAGTGG ACCAGGGGGTT AGCCACCGAC CTTCCGGTCCAC CTTCCGGTCCAC CTTCCGGTCCAC CTTCCGGTCCAC CTTCCGGTCCAC CTTCCGGTCAC CTTCCGGTCACCAC CTTCCGGTCAC CTTCCGCTCACCAC CTTCCGGTCACCAC CTTCCGGTCAC CTTCCGGTCACCAC CTTCCGTCACCA	TGGTCCCCAA ACCAGGGGTT V P N	TCGGTGGCTG AGCCACCGAC R W L	GATGCCAGCC CTACGGTCGG D A S L	TGTCCCCCGT ACAGGGGGCA S P V	CATCCTGGGT GTAGGACCCA I L G	GTCCAGGGTG CAGGTCCCAC V Q G G	GAAGCCAGTG CTTCGGTCAC S Q C	
201	CCTGTCATGT GGACAGTACA L S C	GGGGTGGGGC CCCCACCCCG G V G Q	101 CTIGTCATCT GGGGTGGGGC AGGAGCACTA GAGCCAGTGA ACATCATGGA GCTCTATCTT GGGGCAAGG AATCCAAGAG CTTCACCTTC GGAGGAGAGA CCACGGGTCC TTAGGTTCTC GAAGTGGAAG 68 L S C G V G Q E P T L T L E P V N I M E L Y L G A K E S K S F T F	TCTAACACTA AGATTGTGAT L T L	GAGCCAGTGA CTCGGTCACT E P V N	ACATCATGGA TGTAGTACCT I M E	GCTCTATCTT CGAGATAGAA L Y L	GGTGCCAAGG CCACGGTTCC G A K E	AATCCAAGAG TTAGGTTCTC S K S	CTTCACCTTC GAAGTGGAAG F T F	
101	TACCGGCGGG ATGGCCGCCC Y R R D	ACATGGGGCT TGTACCCCGA M G L	NOT TACCOGCOGO ACATOCCOCA CACOTOCACO CATCOGATICOS CACCOGATICO CACOTOCACO ACOCACACO COATOCACO COATOCACO CACOGATICOS CACOTOCACO CACOTO	TTCGAGTCGG AAGCTCAGCC F E S A	CTGCCTACCC GACGGATGGG A Y P	GGGCTGGTTC CCCGACCAAG G W F	CTGTGCACGG GACACGTGCC L C T V	TGCCTGAAGC ACGGACTTCG P E A	CGATCAGCCT GCTAGTCGGA D Q P	GTCAGACTCA CAGTCTGAGT V R L T	
101	CCCAGCTTCC GGGTCGAAGG	CGAGAATGGT GCTCTTACCA E N G	101 CCCAGCTRCC CGAGAARGOR GCCCCARTOR AGACTFCTAC TECCAGCAGT GTGACTAG GGSTCGAAGG GCCTACACTAC CCGACTTAC GGSTGAAGAT GAGAGAT GAGAGAT AGAGATAGAT 13 OLL P. E. N. G. G. N. N. A. P. I. T. D. P. Y. F. Q. C. D. O.	CCCCCATCAC GGGGGTAGTG P I T	AGACTICTAC TCTGAAGAIG D F Y	TTCCAGCAGT AAGGTCGTCA F Q Q C	GTGACTAG CACTGATC D 0				

CONTIGUACE CONTINUED CAMPAIRAGE RECEIVED CONTROLS RECOGNED RECTURAL METCHAGENS ATCHAGENES TUCCCHARTOS METATORS RECOGNEDAT FORMAN CONTINUED RESONANCE TOWNSHAME CONTINUED RESONANCE TOWNSHAME FORMAN CONTINUED RESONANCE TOWNSHAME FORMAN CONTINUE AND A COLD HAN ON THE GENERAL REPORTS A VALUE OF THE COLD HAND OF THE A CO CACCACCTA COSTICIORAC GOGGICCACA, GAGCCCACACTT CGGICACGA CACTACACC CACCCCGTC TOTA TOTA W L D A S L S P V I L G V Q G S Q C L S C G V G Q E X T L T CHOCHCAR GCCACCTCT CCCCCGTCAT CCTGGGTGTC CAGGGTGGAA GCCAGTGCCT GTCATGTGGG GTGGGGCAGG AGNCGACTCT AACAT 101 201 89

FIG. 8

S	og.	
TAAAATTTC	ATTTTAAAG	
TGTCTTTGCC	ACAGAAACGG	
raggarit teseccticsa geccardart tesecacsas gesacctes titctactra setctearat titccasect tstetisec tarariti	TATCCCTTAA ACCGGGAGCT CCGGTTCTTA AGCCGTGCTC CCCTCGGACG AAAGATGAAT CCAGAGTTTA AAAGGTCGGA ACAGAAACGG ATTT	
GGTCTCAAAT	CCAGAGITIA	
TTTCTACTTA	AAAGATGAAT	
GGGAGCCTGC	GCTC CCCTCGGACG	
TCGGCACGAG	AGCT CCGGTTCTTA AGCCGTGCTC	Aincort ctarte
GGCCAAGAAT	CCGGTTCTTA	Aingo
TGGCCCTCGA	ACCGGGGAGCT	
ATAGGGAATT	TATCCCTTAA	

101 NGCHGTHTAT TTCAAAATAG GGTCTACATA CTGTGAAGCT CATGATGGTT CTGAGTGGGG CACTATGCTT CCGAATGAAG GATTCAGCCT TGAAGGTACT CTAAGTCGGA ACTTCCATGA GGCTTACTTC GTACTACCAA GACTCACCCC GTGATACGAA O orf CCAGATGTAT GACACCTCGA

TGACCTACGG TGAGGAGATC AGTGTTGTCC CAAATCGGGC ACTCCTCTAG TCACAACAGG GTTTAGGCCG V V H H H GRANCHGCAC ANTAACCAGC TGCTGGGTGG AGGACTGCAC GCAGAGAGG TCATTAAAGG CATAGAGGTG TTATTGGTCG AGGACCGACC TCCTGACGTG CGTCTTTCC AGTAATTTCC AEK Ħ G ø 201 GTATCTGCAC AATAACCAGC

GTCGAAGCTT AGGCGACGGA TGGGTCCGAC GAGAAAGGGC CAATTCTGAA ACTTGAGCCA TGAACTCGGT TCCGCTGCCT CTCTTTCCCG GTTAAGACTT CAGCTTCGAA GTCTTACCTC C 301 AGICIGICC CIGICAICCI GGGCGIICAA GGAGGAAGCC AGIGCCIAIC IIGIGGGACA CCTCCTTCGG TCACGGATAG AACACCCTGT CTTCTACCGG CGGGATATGG GCCCTATACC GAAGATGGCC U 401 TGGAGCTCTA CCTCGGGGCC AAGGAATCAA AGAGCTTCAC TCTCGAAGTG 9 9 CCCGCAAGTT TTCCTTAGTT 0 ۵ د GACAGTAGGA GGAGCCCCGG ACCTCGAGAT 53

GATGAAAGTC GATGCTCCCA TCACAGACTT CTACGAGGGT AGTGTCTGAA GINCCICIGO ACCICACOGO AAGCIGACCA GCCIGICAGG CICACICAGA ICCCIGAGGA CCCCGCCIGG GGGGCGGACC P A W CAAGGAGACG TGGAGTGGCC TTCGACTGGT CGGACAGTCC GAGTGAGTCT AGGGACTCCT Н 2 A D S 501

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AGITHMENT ASSECTION SOFTCCCAMA ACTICATIANG CAGAGACIAS GTAGSCAGTG GOSGCTCCTG ATAGAGARTA AGAGACIAGA GTAGACIAGA TCCCCAGAGAC CAGCGAGGAC TATCTCCTAT CACCCTCAC CACCGAGGAC CACCGAGGAC TATCTCCTAT CACCGAGACA 601

AGINGGROS TINCICCITC CCTICCCING IGGACTCCCG CTICTGACCT AAGGCACACA GACACTCTCT ICTCCTGCAT CCCAGIGCTG GTAAATCTTC TCATCCACCG AATGAGGAGA GGAGGGATG ACCTGAGGGC GAAGACTGGA TTCCCGTGTGT CTGTGAGAGA AGAGGACGTA GGGTCACGAC CATTINGAAG 701

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AGAGCAACAT TCTCGTTGTA	CTGGAGAAAG GACCTCTTTC	ATTCTCTGGC TAAGAGACCG	CTGATTCTGG	TATACTTCTC ATATGAAGAG	
BOI ROTHETTOS ACCIOANIST CHARITICIT TOACHITOSA TOGRATIOSA TOGRACIACOS ACCOLATAS ARACCIACOS GRACOACAS AGROCACOS ACCOLACIAS AGROCACOS ACCOLACIAS AGROCACOS TOGRATIASOS TINGGREGAS COTOGRIGAT FUTGORIGAS.	901 AAAGAITOT TOGOTGAAGA AGAGSTGGGA ACTOTTOARA CABAGTAAGA TOTGACACA TACOTGAGAA GACCTGCCBA TOCOTGCGAT TOCOTAGGAT CTGGAGAAAG TTTYCDAAGA ACCACTGCT TCTCCACCCT TGACAAGTAT GFATCATTCT AGACTGTGTC ATGAGATCTT CAGGACGGTA AGGAATACAA GACCTCTTTC	1001 TORMAGESES GENCHAGA CTHTCTCHS CHEGGESC CCTHTCCCTC AACCTITCTS ACATOTSCAG CCTCTCTA TCTTGSCTTC ATCTCTGSC ACCTCCCCC CAGTGSTTCT GAAAGAGAC GACGACCCG GGAAAGAGA TFGGAAAGA TGTAGACGTC GAAGAGATA AGAACGGAG TAAGAGACG	1101 CTERANCEA GAGGGTORIA TCAGRATAGO TGACAGAMA TGACCAGGGA CACTGTOCTG GTTTGAACC AGAGGGGACA ATAAAAAACC CTGATTCTGG GGACTTGGCT CTCCCACTAT AGTCCTATCG ACTGTCTTCT ACTGGTCCGT GTGACAGGAC CAAACTTTGG TCTCCCCTGT TAITTITTGG GACTAGAAC	1201 TOTOTACIO. CATABARGA ROCFFOTBAR CAFTAROTGO GARGAGATTO CTROTAGATA ROMNOCTTO TRAFTORTO TRAFTARAA TATRCTICOT AGRARGAGO GYATTYTOT FOGRACACTT GTRAFTCACO CITOCOTAGO GATGATTART TOTATGGBAC AFTARAGING ARTTARTTA MINIGAGAG	TAGGA
AAACCACGTA TTTGGTGCAT	GTCCTGCCAT	CCTCTCTCAT GGAGAGAGTA	AGAGGGGACA TCTCCCCTGT	TAATTTCATC ATTAAAGTAG	1301 PRINTEREN METTEMAMA ARABADANA ANANARARA ARABADANA ARABADANG GGOGGARG TENTECRIT FROGE AFRANKER TRADANTETE TENTETETE TENTETETE TENTETETE GGOGGGOGG ARABADAR ARABADANER ANANASTRA ANOST ATRADANER TRADANTETET TENTETETETE TENTETETETETETETETETE
AACCCAATAG TTGGGTTATC	TACCTCAGNA ATGGAGTCTT	ACATCTGCAG TGTAGACGTC	GTTTGAAACC CAAACTTTGG	ACATACCTTG	GGCCGCAAGC CCGGCGTTCG
TCTGGTGTGG AGACCACACC	TCTGACACAG AGACTGTGTC	aacciticig tiggaaagac	CACTGTCCTG	CTACTAAATA	AAAAACATGC TTTTTGTACG
TGGTACTACC	CATAGTAAGA	CCTTTCCCTC	TGACCAGGCA	GAAGAGATTG	. Arrararara . Tetetetete
TCAGATTGGA AGTCTAACCT	ACTGTTCATA TGACAAGTAT	CTGGCTGGGC	TGACAGAAGA ACTGTCTTCT	CATTAAGTGG	aaaaaaaaa TTTTTTTT
GTAGATTCTT CATCTAAGAA	AGAGGTGGGA TCTCCACCCT	CTTTCTCTGG	TCAGGATAGC AGTCCTATCG	AGCTTGTGAA TCGAACACTT	AAAAAAAAA TTTTTTTT
AGCTCAATGT TCGAGTTACA	TGGGTGAAGA ACCCACTTCT	GTCACCAAGA CAGTGGTTCT	GAGGGTGATA	CATAAAAAGA GTATTTTTCT	attitaaaa Taaaaittit
TGGTATTTGG ACCATAAACC	aaaagattot ttttctaaga	TGGAGGGGGG	CCTGAACCGA	TCTCTACTCA	TATATTATAT ATATAATATA
801	901	1001	1101	1201	1301

1 GRAGOCIGOT FICTIACITAG GEOTORARIT FECCAGOCTE GEOTFIGOCF ARABITICOS GOTOFFARITA FICHARARIAG GEOTROARIA FECTICAGA COCTOGRACIA DAGAFIGARIC CAGRIGIRAS ACACOTOGAG 101 ATGATGGTTC WGATGGGGC ACTATGCTTC CGAATGAAGG ATTCAGCCTT GAAGGTACTG TATCTGCACA ATAACCAGCT GCTGGCTGGA GGACTGCAG CCTGACGTGC TACTACCAAG ACTCACCCCG TGATACGAAG GCFTACTICC TAAGTCGGAA CTTCCATGAC ATAGACGIGT TATTGGTCGA CGACCGACCT O N N N H T A K V L SAL R M K D CF ы 1 M M V I

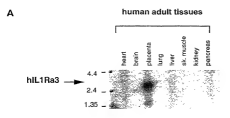
201 CAGAGAAGGT CATTAAAGGT GAGGAGATCA GTGTTGTCCC AAATCGGGCA CTGGATGCCA GTCTGTCCCC TGTCATCCTG GGCGTTCAAG GAGGAAGCCA STOTOTICCA STAATTICCA CICCICIASI CACAACAGGG TITAGCCCGI GACCIACGGI CAGACAGGGG ACAGIAGGAC CCGCAAGIIC CICCITCGGI V I L I S LDAS N R A A A IE D M H E K V

301 GTGCCTATCT TGTGGGACAG AGAAAGGGCC AATTCTGAAA CTTGAGCCAG TGAACATCAT GGAGCTCTAC CTCGGGGCCA AG

CACGGATAGA ACACCTGTC TCTTTCCCGG TTAAGACTTT GAACTCGGTC ACTTGTAGTA CCTCGAGATG GAGCCCGGGT TC 89

hILiRa	1 MEICRGLRSHLITLLLFLFHSETICRPSGRKSS <u>K</u> MQAFRIWDVNQKTFYL
hIL1Rabeta	
tango77	MGSEDWEKDEPOCCLEDPAGSPLEP
hIL1Ra1S	1
hTL1Ra1L	MGSEDWEKDEPQCCLEDPAV SPLEP
hTL1Ra1V	
hIL1Ra2	MRGTPGDADGGG.
hILRa3	
TILLIAN.	
hILIRa	51 B N N
hILlRabeta	25 TINDLNQQVWTLQGQNLVAVPRSDSV
tango77	26 G P
hIL1Ra1S	26 G P
hILlRalL	26 GPSLPAMNEVHTSPKVKNLNPKKESIHDQDHKVLVLDSGNLIAVPDKNYI
hYL1Ra1V	37 GPSLPTMNFVHTSPKVKNLNPKKFSIHDQDHKVLVLDSGNLIAVPDKNYI
hIL1Ra2	13
hILRa3	27 H N N Q L L A G G L H A
IIIIAAS	2 1111
hTL1Ra	S PNVNLEEKIDVVPIEPHALFLGIHGGKMCLSCVKSGDETRLOLEA
hTL1Rabeta	52 TPVTVAVITCKYPEALEGGRODPIYLGIGNPEMCLYCEKVGEO-PTLOL
tango77	36 HIKIFFALASSISSASAE KGSPILLGVSKGEFCLYCDKDKGQSHPSLQL
hTL1Ra1S	36 HTKIFFALASSISSASAE-KGSPILLGVSKGEFCLYCDKDKGQSHPSLQL
hILlRalL	76 RPEIFFALASSLSSASAE - KGSPILLGVSKGEFCLYCDKDKGQSHPSLQL
hIL1RalV	87 RPEIFFALASSLSSASAE-KGSPILLG <u>VSKGEFCLYCDKDKGQSHPSL</u> QL
hIL1Ra2	19 · · IT VAVIT CKYPEAL EQGR GDP IYL GIQN PEMCLYCEKVG EQ · · PT LQL
hILRa3	34 GKVIKGEE ISV VPN RWLDASLSPV ILGVOGGSOCLSCG V G - QEPTLTLEP
hIL1Ra	108 VN IT DL SENRK QD · · KRFAFIR · SDSGPTT SFESAACPGWFLCTAMEADO
hILiRabeta	100 KEQKIMDLYGQPE. PVKPFLFYRAKTGRTSTLESVAFPDWFIASSK-RDQ
tango77	85 KKEKLMKLAAQKESARRPFIFYRAQVGSWNMLESAAHPGWFICTSCNCNE
hIL1Ra1S	85 KKEKL MKLAAQKESARRPFIFYRAQVGSWNMLESAAHPGWFICTSCNCNE
hILlRalL	125 KKEKLMKLAAQKESARRPFIFYRAQVGSWNMLESAAHPGWFICTSCNCNE
hIL1Ra1V	136 KKEKLMKLAAQKESARRPF I FYRAQVGSWNMLE SAAHP GWF I CTSCNCNE
hIL1Ra2	65 KEQKIMDLYGQPE-PVKPFLFYRAKTGRTSTLESVAFPDWFIASSK-RDQ
hILRa3	83 V N I M E L Y L GAKES K S F T F Y R - R D M G L T S S F E S A A Y P G W F L C T V P E A D Q
hIL1Ra	155 PVSLTNMPDEG VMVTKFYFOEDE · · · · · · ·
hIL1Rabeta	148 PIILTSELGKS YNTAFELNIND
tango77	135 PVGVTDKFENR · · · · KHIEFSFQPVCKAEMSPSEVSD
hIL1Ra1S	135 PVGVTDKFENR - · · · KHIEFSFQPVCKAEMSPSEVSD
hillRail	175 PVGVTDKFENR · KHIEFSFQPVCKAEMSPSEVSD
hIL1Ra1V	186 PVGVTDKFENR. · · · KHIEFSFQPVCKAEMSPSEVSD
hIL1Ra2	113 PITT SELGKS YNTAFELNIND · · · · ·
hILRa3	130 PVRLTQLPENGGWNAPITDEYEQOCD





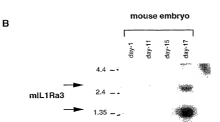
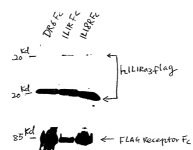


FIG. 11



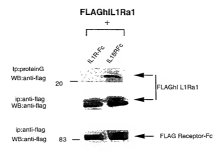


FIG. 13A

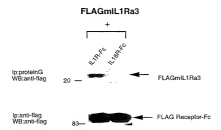


FIG. 14

TCTGGGCCGA CATTCGGGGG ACCTTGGTCC GGGTTCGGAG GGGCGGTACT 1 AAAATGGGCT CTGAGGACTG GGAAAAAGAT GAACCCAGT GCTGCTTAGA AGACCCGGCT GTAAGCCCCC TGGAACCAGG CCCAAGCCTC CCGGCCATGA м Ø > Δ, Ω TITIACCCGA GACTCCTGAC CCTTITTCTA CTTGGGGTCA CGACGAATCT U Д ы Ω × ы Д ^insert starts

101 ATTITOTICA CACAAGICCA AAGGIGAAGA ACTITAAACCC GAAGAAATIC AGCATICATG ACCAGGAICA CAAAGTACIG GICCIGGACI CIGGGAATCI TGAAFITIGGG CITCTTTAAG TCGTAAGTAC IGGTCCTAGT GTTTCATGAC CAGGACCTGA GACCCTTAGA L > × н α SIHD tr. × L N z GIGITCAGGI IICCACIICI K V K TAAAACAAGT

CGGAGACGCC TCTTTCCTTC AGGCTAAGAG 201 CAUNGCAGTT CCAGADAAAA ACTACATACG CCCAGAGATC TTCTTTGCAT TAGCCTCATC CTTGAGCTCA GCCTCTGCGG AGAAAGGAAG A A S GHALCGICAA GGICTATITI IGAIGIAIGC GGGICICIAG AAGAAACGIA AICGGAGTAG GAACTCGAGI S ü A F HEL YIR PDKN N V 67 301 CTGGGGGTCT CTAAAGGGGA GTTTTGTCTC TACTGTGACA AGGATAAAGG ACAAAGTCAT CCATCCCTTC AGCTGAAGAA GGAGAAACTG ATGAAGCTGG CCTCTTTGAC TCGACTTCTT L K TGTTTCAGTA GGTAGGGAAG S S α ATGACACTGT TCCTATTTCC × D υ × GACCCCCAGA GATTTCCCCT CAAACAGAG [z,

CGAGTGGGGC GGAGTCGGCG CCTCAGCCGC s [1] CGCCGGCCCT TCATCTITIA TAGGGCTCAG GTGGGCTCCT GGAACATGCT CCTTAGTCGT GCGGCCGGGA AGTAGAAAT ATCCCGAGTC CACCCGAGGA CCTTGTACGA N z S o A α R CTGCCCAAAA GGAATCAGCA SA ы GACGGGTTTT 40.1 134 TGGGGTGACA GATAAATTTG AGAACAGGAA ACACATTGAA TTTTCATTTC AACCAGTTTG CAAAGCTGAA ACGTTAACAT TACTCGGACA ACCCCACTGT CTATTTAAAC TCTTGTCCTT TGTGTAACTT AAAAGTAAAG TTGGTCAAAC GTTTGGACTT E) Ξ K > ပ TGCAATTGTA ATGAGCCTGT ь GACGTGGAGG 501

601 ATGAGCCCCA GTGAGGTCAG CGATTA TACTCGGGGT CACTCCAGTC GCTAAT

200 M S P S E V S

C CCCGCCATGA G GGGCGGTACT P A M N
TGGAACCAGG CCCAAGCCTC CCCGCCATA ACCTTGGTCC GGGTTCGGAG GGGCGGTAA E P G P S L P A M
TGGAACCAGG ACCTTGGTCC E P G
AAATGGSCT CTGAGGACTG GGAAAAAAA GAA GAACCCGGC GGAAACCAGG CCCAAGCCTC TTTACCCGAA GACTCCTGA CCTTTACCCGAA GACTCGGAG CCTTTACCCGAA GACTCGGAG CCTTTACCTCAA CTTGGGGGCA CATTCGGAGG ACTTTGGATCC GGGGTTCGGAG A S E B W E K D E P Q C C L E D P A V S P L E P G P S L insert starts
AGACCCGGCT G' TCTGGGCCGA CJ D P A V
AF GAACCCCAGT GCTGCTTAGA AGACCO
GAACCCCAGT CTTGGGGTCA E P Q C
GGAAAAGAT CCTTTTTCTA E K D
GGGCT CTGAGGACTG GGAAAAAGAT CCCGA GACTCCTGAC CCTTTTTTTA G S E D W E K D Ft starts
1 AAAATGGGCT CTGAC TTTTACCCGA GACTC 1 M G S E ^insert starts ^orf
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101 ATTIVITCA CACAAGAIC TICTTICCAT TAGCCICATC CTICAGCICA GCTICAGCGA AGAAAGAGGAA TAAAACAAG GTGTTICTAG AAGAAACGTA ATCGGAGTAGGAGGC GCTTTICCTIC AGGCTAAGAG GACCCCCAAA GATTICCCCT 34 P V H T K I F P A L A S S L S A S A E K G S P I L L G V S K G E	201 GTTTTGTCTC TACTGTGA.CA AGGAINAAGGA ACANAGTCAT CCATCCCTTC AGCTGAAGAA GGAGGAACTG ATGGAGCTGG CTGCCCAAAA GGANTCAGCA CAAAACAGAG ATGACACTGT TCCTATTTCC TGTTTCAGTA GGTRAGGAAG TCGACTTCTT CCTCTTTGAC TACTTCGAC GACGGGTTTT CCTTAGTCGT 67 F C L Y C D K D K G Q S H P S L Q L K K E K L M K L A A O K F S A
AAA TTT	AAT TTA
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0000	366.
CTG	CTG
CTC	ACC ACC
TAA	AGC. TCG2
2000 MGGC	ACT ACT
AG J	7 D
CCT	AAC
SAAA CTTT	SAGA
S C B	85.
9000 V	AGA TCT
SAGA	SACT N K
90 A	AGC TCC
SAGT	AAAG
GAGC	AGGG
CTT GAA L	CCA
ATC IAG S	STA
CTC GAG	AGT TCA(
TAGC ATCG	ACAA IGITT
AT I	9000
LACG AACG	PAAA VITT
AGA?	SGAT CCTA
E A F	A T A
AGAT ICTA	rgac ACTG D
GTT	GAC
2 P	AT S
AAG; H	TCTC AGAC
AAC V V	AAC
ATA AAT	CA.
101	201

301 CGCCGGCCCT TCATCTTTA TAGGGCTCAG GTGGCCTCCT GGAACATGCT GGAGTCGCCG GCCCCCC GATGGTTCAT CTGCACCTCC TGCAATTGTA	401 ATGAGCCTGT TGGGGGGACA GATAAATTG AGAACAGGAA ACACATTGA TTTCATTTC AACCAGTTTG CAAGCTGAA ATGAGCCCCA GTGAGGTCAG
GCGGCCGGGA AGTAGAAAA ATCCCGAGTC CACCCGAGGA CCTTGTAGAG CCTCAGCCGC CGAGTGGGGG CTACCAAGTA GACGTGGAGG ACGTTAACAT	TACTCGGACA ACCCACTGT CTATTTAAAC TGTTGTCTT TGTGTAACTT AAAAGTAAG TTGGTCAAAC GTTTCGACTT TACTCGGGGT CACTCAGC
100 R R P I F Y R A Q V G S W N M L E S A A H P G W F I C T S C N C N	134 E P V G V T D K P E N R K H I E F S F Q P V C K A E M S P S E V S
CTGCACCTCC	ATGAGCCCCA
GACGTGGAGG	TACTCGGGGT
C T S	M S P S
GATGGTTCAT	CAAAGCTGAA
CTACCAAGTA	GTTTCGACTT
W F I	K A E
GCTCACCCCG	AACCAGTITG
CGAGTGGGGC	TTGGTCAAAC
A H P G	P V C
GGAGTCGGCG	TTTTCATTTC
CCTCAGCCGC	AAAAGTAAAG
E S A	FSFQ
GGAACATGCT	ACACATTGAA
CCTTGTACGA	TGTGTAACTT
N M L	H I E
GTGGGCTCCT	AGAACAGGAA
CACCCGAGGA	TCTTGTCCTT
V G S W	N R K
TAGGGCTCAG	GATAAATTTG
ATCCCGAGTC	CTATTTAAAC
R A Q	D K P E
TCATCTTTA	TGGGGTGACA
AGTAGAAAAT	ACCCCACTGT
I F Y	G V T
CGCCGGCCCT	401 ATGAGGCTGT TGGGGTGAG GATAAATTTG AGAACAGGAA ACACATTGAA TTTTCATTTC AACAGTTTG CAAAGGTGAA ATGAGGCCCA GTGAGGTCAG
GCGGCCGGGA	TACTCGGACA ACCCACTGT TCTTTTAAAC TCTTGTCTT TGTGTAACTT AAAAGTAAAG TTGGTCAAAC GTTTCGACTT TACTCGGGGT CACTCCAGTC
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1 AACCCGAAGA AATTCAGCAT TCATGACCAG GATCACAAAG TACTGGTCCT GGACTCTGGG AATCTCATAG CAGTTCCAGA TAAAAACTAC ATACGCCCAG TTAGAGTATC GTCAAGGTCT ATTTTTGATG TATGCGGGTC > TIGGGCTICT TIAAGICGIA AGIACIGGIC CIAGIGITIC AIGACCAGGA CCIGAGACCC Д r v ٥ DHK 0 П

TCAICCTIGA GCICAGCCIC IGCGGAGAAA GGAAGICCGA IICICCIGGG GGICICIAAA GGGGAGIIII GICICIACIG CCCCTCAAAA CAGAGATGAC TCTAGAAGAA ACGTAATCGG AGTAGGAACT CGAGTCGGAG ACGCCTCTTT CCTTCAGGCT AAGAGGACCC CCAGAGATTT V S K GSPI A S 101 AGATCTTCTT TGCATTAGCC

TONCANGOAT AAAGGACAAA GTCATCCATC CCTTCAGCTG AAGAAGGAGA AACTGATGAA GCTGGCTGCC CAAAAGGAAT CAGCAGGCG GCCCTTCATC GTTTTCCTTA GTCGTGCGGC CGGGAAGTAG O K E CGACCGACGG L A A ACTGTTCCTA TYTCCTGTTT CAGTAGGTAG GGAAGTCGAC TTCTTCCTCT TTGACTACTT K K E K 7 0 П E P 201

TIGIAATGAG CCTGTIGGGG AACATTACTC CCTCCTGCAA GGGGCCTACC AAGTAGACGT GGAGGACGTT E U 301 TITIATAGG CTCAGGTGG CTCCTGGAAC ATGCTGGAGT CGGCGCTCA CCCCGGATGG TTCATCTGCA M O Δ, AAAATATCCC GAGTCCACCC GAGGACCTTG TACGACCTCA GCCGCCGAGT A ď M N > 0 101

CAGTCGCTAA TCCTTTGACG 401 TGACAGATAA ATTIGAGAAC AGGAAACACA TIGAATITIC ATTICAACCA GITIGCAAAG CIGAAAIGAG CCCCAGIGAG GICAGGGAIT AGGAAACIGC GGGGTCACTC S ۵ ACTGICTAIT TAAACTCTIG TCCTTIGIGT AACTTAAAAG TAAAGTIGGT CAAACGITIC GACTTIACTC Ø z ы O A ۵ o ſz, ET. × × z u Б., 135

501 CCCATTGAAC GCCTTCCTCG CTAATTTGAA CTAATTGTAT AAAAACACCA AACCTGCTCA SGGTAACTTG CGGAAGGAGC GATTAAACTT GATTAACATA TTTTTGTGGT TTGGACGAGT

hILlRalL		]
hIL1Ra1V	1 MSFVGENSGVKMGSEDWEKDEPQCCLEDPAGSPLEPGPSLPTMNFVHTSP	
hIL1Ra1S	1 ·····	j
hIL1Ra1	1 - V K N L N P K K F S I H D Q D B K V L V L D S G N - · · · · · · · · · · · · · · · · · ·	
hILlRalL	2 GSEDWEKDEP OCCLEDPAV SPLEP GP SLPAMN FV HT SPK V K N L N P K K F S I	
hIL1Ra1V	51 K V K N L N P K K F S I H D O D H K V L V L D S G N - · · · · · · · · · · · · · · · · · ·	
hILlRa1S	Z GSEDWEK DEPOCCLED PAVSPLEPGP - · · · · · · · · · · · · · · · · · ·	
hIL1Ra1	26 LIAVPDKNYIRPEIFFALASSLSSASAEKGSPILLG	1
hILlRalL	52 HOODHKVLVLDSGNLIAVPDKNYIRPEIFFALASSLSSASAEKGSPILLG	1
hIL1Ra1V	TT L T A V P D K N Y I R P E I F F A L A S S L S S A S A E K G S P I L L G	1
hIL1Ra1S	28 SLPAMNFVHTKIFFALASSLSSASAEKGSPILLG	
		•
hIL1Ra1	62 VSKGEFCLYCDKDKGQSHPSLQLKKEKLMKLAAQKESARRPFIFYRAQVG	1
hILlRalL	102 VSKGEFCLYCDKDKGQSHPSLQLKKEKLMKLAAQKESARRPFIFYRAQVG	ı
hIL1Ra1V	113 VSKGEFCLYCDKDKGQSHPSLQLKKEKLMKLAAQKESARRPFIFYRAQVG	
hIL1Ra1S	62 V S K G E F C L Y C D K D K G Q S H P S L Q L K K E K L M K L A A Q K E S A R R P F I F Y R A Q V G	
		•
hIL1Ra1	122 SWNMLESAAHPGWFICTSCNCNEPVGVTDKFENRKHIEFSFQPVCKAEMS	1
hIL1Ra1L	152 SWNMLESAAHPGWFICTSCNCNEPVGVTDKFENRKHIEFSFQPVCKAEMS	l
hIL1Ra1V	163 SWN ML E SAAHPGWF I CT SCN CN EP V G V T D K F E N R K H I E F S F Q P V C K A E M S	l
hIL1Ra1S	112 SWNMLESAAHPGWFICTSCNCNEPVGVTDKFENRKHIEFSFQPVCKAEMS	1
		•
hILlRa1	162 PSEVSD	
hILlRalL	202 P S E V S D	
hTL1Ra1V	213 P S E V S D	
hILLRalS	362 P S E V S D	

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- GGCCCAAGCC TCCCCACCAT CCGGGTTCGG AGGGGTGGTA Ö Δ CTCTGAGGAC TGGGAAAAG ATGAACCCCA GTGCTGCTTA GAAGACCCGG CTGGAAGCCC CCTGGAACCA GAGACTCCTG ACCCTTTTTC TACTTGGGGT CACGACGAAT CTTCTGGGCC GACCTTCGGG GGACCTTGGT D: 101 TGAAAATGGG ACTITIACCC
  - TGACCAGGAT CACAAAGTAC TGGTCCTGGA GCCTTCTTTA AGTCGTAAGT ACTGGTCCTA GTGTTTCATG ACCAGGACCT a U e e ... ro. Ö CCGAAGAAT TCAGCATTCA ρ. E υ GAACTTAAAC CTTGAATTTG e. WEKD CAAAGGTGAA GTTTCCACTT CACACAAGTC GTGTGTTCAG c GAATTTTGTT CTTAAAACAA 201 11
    - t) GGAGAAAGGA ... > V L TCCTTGAGCT CAGCCTCTGC × 0 Д TCTTCTTTGC ATTAGCCTCA TAATCGGAGT SIH PKKF GCGGGTCTCT AGAAGAACG CTCATAGCAG TTCCAGATAA AAACTACATA CGCCCAGAGA 2 N TTTGATGTAT × К GAGTATCGTC AAGGTCTATT r) Ĺ 44 301
      - CCTCTTTCCT ы AGGAACTCGA GTCGGAGACG A S A SIS V) L A F F RPET N X I z D Д LIAV 77
- TICCTCTITG ACTACTICGA CTCTAAAGGG GAGTTTTGTC TCTACTGTGA CAAGGATAAA GGACAAAGTC ATCCATCCCT TCAGCTGAAG AAGGAGAAAC AGTCGACTTC O L K CTCAAAACAG AGATGACACT GTTCCTATTT CCTGTTTCAG TAGGTAGGGA I S d G Q S H K D K O O × C EL, GAGATTTCCC Ü S TCCTGGGGGT AGGACCCCCA 401 111
- AAGGAATCAG CACGCCGGCC CTTCATCTTT TATAGGGCTC AGGTGGGCTC CTGGAACATG CTGGAGTCGG CGGCTCACCC GTGCGGCCGG GAAGTAGAAA ATATCCCGAG TCCACCCGAG GACCTTGTAC GACCTCAGCC S ... E11 N O > Ø YRA <u>п</u> ſı. œ æ TICCITAGIC EXI GGCTGCCCAA CCGACGGGTT K 501 144
- GTTGGGGTGA CAGATAAATT TGAGAACAGG AAACACATTG AATTTTCATT TCAACCAGTT TTAAAAGTAA AGTTGGTCAA TTTGTGTAAC GTCTATTTAA ACTCTTGTCC Ω GGACGTTAAC ATTACTCGGA CAACCCCACT > O Ν CCTGCAATTG TAATGAGCCT м z z ATCTGCACCT TAGACGTGGA 177 I C 601
- ATTTGAACTA ATTGTATAAA AACACCAAAC CTGCTCACTA TITGACGGGG TAACTTGCGG AAGGAGCGAT TAAACTTGAT TAACATATTT TIGTGGTTTG GACGAGTGAT AAACTGCCCC ATTGAACGCC TTCCTCGCTA CAGTGAGGTC AGCGATTAGG TCGCTAATCC SD GTCACTCCAG SEV AAATGAGCCC TTTACTCGGG 701 211
- AAAAAAAAA AAAAAACGT TTGCGGCCGC AAGCTTATT ITTITITIT TITITIGCA AACGCCGGCG TICGAATAA 801

### COMBINED DECLARATION FOR PATENT APPLICATION AND POWER OF ATTORNEY

As a below named inventor, I hereby declare that:

the specification of which (check only one item below): is attached hereto.

2001).

9555

and was amended on \_\_\_\_\_

the claims, as amended by any amendment referred to above.

My residence, post office and citizenship are as stated below next to my name,

was filed as United States application Serial No.

I believe I am the original and sole inventor (if only one name is listed below) or an original, first and joint inventor (if plural names are listed below) of the subject matter which is claimed and for which a patent is sought on the invention entitled:

#### IL-1 RELATED POLYPEPTIDES

was amended under PCT Article 19 on ( ) (if applicable) (now assigned as U.S.S.N 09/869,566, filed June 29,

Harby state that I have reviewed and understand the contents of the above-identified specification, including

was filed as PCT international application Number PCT/US99/30720 on 22 December 1999 and

acknowledge the du accordance with Title (	ity to disclose information which 37, Code of Federal Regulations,	ch is material to the examina §1.56(a).	ation of this application in
patent or inventor's or han the United States or inventor's certificat United States of Ame opplication(s) of which	APPLICATION/U.S. PROVISIONA	onal application(s) designating a also identified below any fore cation(s) designating at least of subject matter having a filin	at least one country other ign application(s) for patent one country other than the g date before that of the
COUNTRY IS IS IS	APPLICATION NUMBER 60/113,430 60/116,843 60/129,122	DATE OF FILING (day, month, year) 23 December 1998 22 January 1999 13 April 1999	PRIORITY CLAIMED UNDER 35 USC 119

I hereby claim the benefit under Title 35, United States Code, §120 of any United States application(s) or PCT international application(s) designating the United States of America that is/are listed below and, insofar as the subject matter of such of the claims of this application is not disclosed in that/those priori application(s) in the manner provided by the first paragraph of Title 35, United States Code §112, I acknowledge the duty to disclose material information a defined in Title 37, Code of Federal Regulations, §1.56(a) which occurred between the filing date of the prior application(s) and the national or PCT international filing date of this application:

# COMBINED DECLARATION FOR PATENT APPLICATION AND POWER OF ATTORNEY (Continued) PRIOR U.S. APPLICATIONS OR PCT INTERNATIONAL APPLICATIONS OF PCT INTERNATIONAL APPLICATION A

THIO	THE U.S. FOR	OR PCT INTERNATIONAL R BENEFIT UNDER 35 U.S	APPLICATIONS I.C. 120:	DESIGNATIN	G
U.S. APPLICATIONS		STATUS (check one)			
U.S. Application Num	ber	U.S. Filing Date	Patented	Pending	Abandoned
PCT APPLICATIONS [	DESIGNATING THE U.S.				
PCT Application No.	PCT Filing Date	U.S. Serial Numbers			
PCT/US99/30720	22 December 1999				
Send correspondence to	Genentech, Inc. Attn: Craig G. 1 DNA Way South San Franc Telephone: (66	Svoboda cisco, CA 94080-4990 0) 225-1489	demark Office con	nected therewith	
or both, under Section 1 application or any patent The undersigne his foreign pate without direct o	001 of Title 18 of the Unite t issue thereon.  d hereby authorizes the U.S and agent as to any action to communication between the m whom instructions may be	f my own knowledge and bel ful false statements and the li d States Code and that willful . attorney or agent named her be taken in the Patent and T U.S. attorney or agent and the e taken, the U.S. attorney or	false statements	may jeopardize t	the validity of the
Full name of sole or first	inventor				<del></del>
Audrey Goddard					
nventor's Name		Inventor's signature		D	ate
I 10 Congo Street, San F Residence	rancisco, California 94131				
Canada					
Citizenship					
DNA Way, South San F	rancisco, CA 94080-4990				
ost Office Address					

Page 2 of 3

Full name of second joint inventor, if an		<b>-</b> . /
Guohua James Pan		Feb 11/2002
Inventor's Name	Inventor's signature	Date
14 Tewsley Place, Etobicoke, Ontario, (	Canada, M9P 1N7	
Residence		
Canada		
Citizenship		
1 DNA Way, South San Francisco, CA S	94080-4990	

Post Office Address

### COMBINED DECLARATION FOR PATENT APPLICATION AND POWER OF ATTORNEY

As a below named inventor, I hereby declare that:

My residence, post office and citizenship are as stated below next to my name,

I believe I am the original and sole inventor (if only one name is listed below) or an original, first and joint inventor (if plural names are listed below) of the subject matter which is claimed and for which a patent is sought on the invention entitled:

	IL-1 RELATED	POLYPEPTIDES	
the specificati	on of which (check only one item below):		
_	is attached hereto.		
_	was filed as United States application Se and was amended on	erial No on	
2001).	was filed as PCT international application under PCT Article 19 on ( ) (if applicable)	e) (now assigned as U.S.S.N (	on 22 December 1999 and 09/869,566, filed June 29,
hereby state the claims, as	that I have reviewed and understand the amended by any amendment referred to a	e contents of the above-identi bove.	
acknowledge accordance w	e the duty to disclose information which th Title 37, Code of Federal Regulations,	th is material to the examina §1.56(a).	ation of this application in
than the Unite for inventor's a United States application(s)	foreign priority benefits under Title 35, ntor's certificate or of any PCT internati d States of America listed below and have certificate or any PCT international applic of America filed by me on the same of which priority is claimed.	onal application(s) designating e also identified below any fore ation(s) designating at least c subject matter having a filing	at least one country other ign application(s) for patent one country other than the g date before that of the
TO PRIOR FORE	IGN/PCT APPLICATION/U.S. PROVISIONA U.S.C	L APPLICATIONS WHICH CLA C. 119:	IMS PRIORITY UNDER 35
COUNTR US US US	Y APPLICATION NUMBER 60/113,430 60/116,843 60/129,122	DATE OF FILING (day, month, year) 23 December 1998 22 January 1999 13 April 1999	PRIORITY CLAIMED UNDER 35 USC 119
international a	the benefit under Title 35, United States oplication(s) designating the United States of such of the claims of this application	of America that is/are listed	below and, insofar as the

manner provided by the first paragraph of Title 35, United States Code §112, I acknowledge the duty to disclose material information a defined in Title 37, Code of Federal Regulations, §1.56(a) which occurred between the filing date of the prior application(s) and the national or PCT international filing date of this application:

## COMBINED DECLARATION FOR PATENT APPLICATION AND POWER OF ATTORNEY (Continued) PRIOR U.S. APPLICATIONS OR PCT INTERNATIONAL APPLICATIONS DESIGNATING

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Post Office Address

U.S. APPLICATIONS  U.S. Application Number  U.S. Filing Date  Patented  Pending  Abandone  Per Application Number  Per Application No.  Per Filing Date  Per Application No.  Per Filing Date  Per Application No.  Per Filing Date  U.S. Serial Numbers  Per Application No.  Per Filing Date  Power Of ATTORNEY: As a named inventor, I hereby appoint the following attorney(s) and/or agent(s) to prosecute this application and transact all business in the Patent and Trademark Office connected therewith.  Description of Attr. Craig G. Svoboda  I DNA Way  South San Francisco, CA 94080-4990  Telephone: (650) 225-1489  Per bety declare that all statements made herein of my own knowledge and belief are believed to be true; and further that is statements were made with the knowledge that willful false statements and the like so made are punishable by fine or imprison or both, under Section 1001 of Title 18 of the United States Code and that willful false statements may be stated the validity of application or any patent issue thereon.  The undersigned hereby suthorizes the U.S. attorney or agent named herein at accept and follow instructions from his foreign patent agent as to any action to be taken in the Patent and Trademark Office regarding this application without direct communication between the U.S. attorney or agent named herein at accept and follow instructions may be taken, the U.S. attorney or agent named herein will be so notified by the undersigned.  Per Pending Abandone  Altered Pending Abandone  Per Attrace Residence  Per Attrace Residence  Per Attrace Residence  Per Attrace Residence  Pending Abandone  Per Attrace Residence  Pending Abandone  Per Attrace Residence  Pending Abandone  Pending A	PRIO	R U.S. APPLICATIONS C THE U.S. FOR	R PCT INTERNATIONAL A BENEFIT UNDER 35 U.S.	APPLICATIONS .C. 120:	DESIGNATIN	G ,
PCT APPLICATIONS DESIGNATING THE U.S.  PCT Application No. PCT Filing Date U.S. Serial Numbers  PCT/US99/30720 22 December 1999  POWER OF ATTORNEY: As a named inventor, I hereby appoint the following attorney(s) and/or agent(s) to prosecute this application and transact all business in the Patent and Trademark Office connected therewith.  PCT Application on Attr: Craig G. Svoboda  1 DNA Way South San Francisco, CA 94080-4990 Telephone: (650) 225-1489  Pereby declare that all statements made herein of my own knowledge and belief are believed to be true; and further that the statements were made with the knowledge that willful false statements and the like so made are punishable by fine or imprison or both, under Section 1001 of Title 18 of the United States Code and that willful false statements may jeopardize the validity of application or any patent issue thereon.  The undersigned hereby authorizes the U.S. attorney or agent named herein at accept and follow instructions from his foreign patent agent as to any action to be taken in the Patent and Trademark Office regarding this application without direct communication between the U.S. attorney or agent and the undersigned. In the vent of a change in the persons from whom instructions may be taken, the U.S. attorney or agent named herein will be so notified by the undersigned.  PCL ADMAND INVENTOR SNAME INVENTOR SNAME Inventor's signature Date  Date  Canada			STATUS (check one)			
PCT Application No. PCT Filing Date U.S. Serial Numbers  PCT/US99/30720 22 December 1999  POWER OF ATTORNEY: As a named inventor, I hereby appoint the following attorney(s) and/or agent(s) to prosecute this application and transact all business in the Petent and Trademark Office connected therewith.  O9157  LIENT TRADEMARK.OPPICE  Send correspondence to Genentach, Inc.  Attr: Craig G. Svoboda  1 DNA Way  South San Francisco, CA 94080-4990  Telephone: (650) 225-1489  Thereby declare that all statements made herein of my own knowledge and belief are believed to be true; and further that of statements were made with the knowledge that willful false statements and the like so made are punishable by fine or imprison or both, under Section 1001 of Tritle 18 of the United States Code and that willful false statements may jeopardize the validity of application or any patient issue thereon.  The undersigned hereby authorizes the U.S. attorney or agent named herein at accept and follow instructions from his foreign patient agent as to any action to be taken in the Patient and Trademark Office regarding this application without direct communication between the U.S. attorney or agent and the undersigned. In the vent of a change in the patients of my whom instructions may be taken, the U.S. attorney or agent named herein will be so notified by the undersigned.  Full name of sole or first inventor  Modrey Goddard  Inventor's Name  Inventor's Name  Inventor's Name  Little Congo Street, San Francisco, Celifornia 94131  Afficiency Date of Trademark Office connected therewith.	U.S. Application Num	ber	U.S. Filing Date	Patented	Pending	Abandoned
PCT Application No. PCT Filing Date U.S. Serial Numbers  PCT/US99/30720 22 December 1999  POWER OF ATTORNEY: As a named inventor, I hereby appoint the following attorney(s) and/or agent(s) to prosecute this application and transact all business in the Petent and Trademark Office connected therewith.  O9157  LIENT TRADEMARK.OPPICE  Send correspondence to Genentach, Inc.  Attr: Craig G. Svoboda  1 DNA Way  South San Francisco, CA 94080-4990  Telephone: (650) 225-1489  Thereby declare that all statements made herein of my own knowledge and belief are believed to be true; and further that of statements were made with the knowledge that willful false statements and the like so made are punishable by fine or imprison or both, under Section 1001 of Tritle 18 of the United States Code and that willful false statements may jeopardize the validity of application or any patient issue thereon.  The undersigned hereby authorizes the U.S. attorney or agent named herein at accept and follow instructions from his foreign patient agent as to any action to be taken in the Patient and Trademark Office regarding this application without direct communication between the U.S. attorney or agent and the undersigned. In the vent of a change in the patients of my whom instructions may be taken, the U.S. attorney or agent named herein will be so notified by the undersigned.  Full name of sole or first inventor  Modrey Goddard  Inventor's Name  Inventor's Name  Inventor's Name  Little Congo Street, San Francisco, Celifornia 94131  Afficiency Date of Trademark Office connected therewith.						
PCT Application No. PCT Filing Date U.S. Serial Numbers  PCT/US99/30720 22 December 1999  POWER OF ATTORNEY: As a named inventor, I hereby appoint the following attorney(s) and/or agent(s) to prosecute this application and transact all business in the Patent and Trademark Office connected therewith.  POPIES TRADEMARS.OPECS  Send correspondence to  Genentech, Inc.  Attr: Craig G. Svoboda  1 DNA Way  South San Francisco, CA 94080-4990  Telephone: (650) 225-1489  Persety declare that all statements made herein of my own knowledge and belief are believed to be true; and further that of statements were made with the knowledge that willful false statements and the like so made are punishable by fine or imprison or both, under Section 1001 of Tritle 18 of the United States Code and that willful false statements may jeopardize the validity of supplication or any patent issue thereon.  The undersigned hereby authorizes the U.S. attorney or agent named herein at accept and follow instructions from his foreign patent agent as to any action to be taken in the Patent and Trademark Office regarding this application without direct communication between the U.S. attorney or agent and the undersigned. In the vent of a change in the persons from whom instructions may be taken, the U.S. attorney or agent named herein will be so notified by the undersigned.  Polyton of the property of the property of the U.S. attorney or agent named herein will be so notified by the undersigned.  Polyton of the property of the U.S. attorney or agent named herein will be so notified by the undersigned.  Polyton of the U.S. attorney or agent named herein will be so notified by the undersigned.  Polyton of the U.S. attorney or agent named herein will be so notified by the undersigned.						
PCT Application No. PCT Filing Date U.S. Serial Numbers  PCT/US99/30720 22 December 1999  POWER OF ATTORNEY: As a named inventor, I hereby appoint the following attorney(s) and/or agent(s) to prosecute this application and transact all business in the Patent and Trademark Office connected therewith.  POPIES TRADEMARS.OPECS  Send correspondence to  Genentech, Inc.  Attr: Craig G. Svoboda  1 DNA Way  South San Francisco, CA 94080-4990  Telephone: (650) 225-1489  Persety declare that all statements made herein of my own knowledge and belief are believed to be true; and further that of statements were made with the knowledge that willful false statements and the like so made are punishable by fine or imprison or both, under Section 1001 of Tritle 18 of the United States Code and that willful false statements may jeopardize the validity of supplication or any patent issue thereon.  The undersigned hereby authorizes the U.S. attorney or agent named herein at accept and follow instructions from his foreign patent agent as to any action to be taken in the Patent and Trademark Office regarding this application without direct communication between the U.S. attorney or agent and the undersigned. In the vent of a change in the persons from whom instructions may be taken, the U.S. attorney or agent named herein will be so notified by the undersigned.  Polyton of the property of the property of the U.S. attorney or agent named herein will be so notified by the undersigned.  Polyton of the property of the U.S. attorney or agent named herein will be so notified by the undersigned.  Polyton of the U.S. attorney or agent named herein will be so notified by the undersigned.  Polyton of the U.S. attorney or agent named herein will be so notified by the undersigned.	PCT APPLICATIONS O	DESIGNATING THE II C				
POWER OF ATTORNEY: As a named inventor, I hereby appoint the following attorney(s) and/or agent(s) to prosecute this application and transact all business in the Patent and Trademark Office connected therewith.    O9157			II C Carial Numbers			
Prosecute this application and transact all business in the Patent and Trademark Office connected therewith.  915.7  NENT TRADEMARK OFFICE  Send correspondence to  Genentech, Inc.  Attn: Craig G. Svoboda 1 DNA Way South San Francisco, CA 94080-4990 Telephone: (650) 225-1489  Intereby declare that all statements made herein of my own knowledge and belief are believed to be true; and further that the statements were made with the knowledge that willful false statements and the like so made are punishable by fine or imprison or both, under Section 100.1 of Title 18 of the United States Code and that willful false statements may jeporadize the validity of application or any patent issue thereon.  The undersigned hereby authorizes the U.S. attorney or agent named herein at accept and follow instructions from his foreign patent agent as to any action to be taken in the Patent and Trademark Office regarding this application without direct communication between the U.S. attorney or agent and the undersigned. In the event of a change in the persons from whom instructions may be taken, the U.S. attorney or agent named herein will be so notified by the undersigned.  Subject to the properties of the properties of the patent and the undersigned. In the event of a change in the persons from whom instructions may be taken, the U.S. attorney or agent named herein will be so notified by the undersigned.  Subject to the properties of the patent and Trademark Office regarding this application without direct communication between the U.S. attorney or agent and the undersigned. In the event of a change in the persons from whom instructions may be taken, the U.S. attorney or agent named herein will be so notified by the undersigned.		-	U.S. Serial Numbers			
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pr both, under Section 1001 of Title 18 of the United States Code and that willful false statements may jeopardize the validity of application or any patent issue thereon.  The undersigned hereby authorizes the U.S. attorney or agent named herein at accept and follow instructions from his foreign patent agent as to any action to be taken in the Patent and Trademark Office regarding this application without direct communication between the U.S. attorney or agent and the undersigned. In the rest of a change in the persons from whom instructions may be taken, the U.S. attorney or agent named herein will be so notified by the undersigned.  Full name of sole or first inventor  Audrey Goddard  Date  110 Congo Street, San Francisco, California 94131  LA  Canada  Canada  Canada  Canada  Canada  Canada  Canada	prosecute this	Genentech, Inc. Attn: Craig G. S 1 DNA Way South San Franc Telephone: (656	09157  O9157  O9157  O9157  O9157  O9157  O9157  O9167  O9	lief are believed i	nected therewith	further that these
Audrey Goddard  O: GodWard  Inventor's Name  Inventor's signature  Date  110 Congo Street, San Francisco, California 94131  Canada	or both, under Section 1 application or any paten  The undersigne his foreign pate without direct of the persons fro	001 of Title 18 of the Unite t issue thereon.  Id hereby authorizes the U.S ant agent as to any action to communication between the m whom instructions may be	d States Code and that willful  attorney or agent named her be taken in the Patent and T U.S. attorney or agent and the	I false statements rein at accept and Frademark Office e undersigned. In	may jeopardize  I follow instructi regarding this ap	the validity of the ons from oplication
110 Congo Street, San Francisco, California 94131  Canada	Audrey Goddard	inventor	D: Soddard			
		rancisco, California 94131	LA			
o uzonomp						
I DNA Way, South San Francisco, CA 94080-4990	·					

	Docket No.	P2534-3
4	2.1	4

Foll name of second joint inventor, in	anv	
2.60	,	
Guohua James Pan		
Inventor's Name	Inventor's signature	Date
	4 10 /	
14 Tewsley Place, Etobicoke, Ontari	o, Canada, M9P 1N7 // # //	
Residence		
Canada		
Citizenship		
·		
1 DNA Way, South San Francisco, C	A 94080-4990	
Post Office Address		

## RECONSTATIO 19 FEB 2002

# #5

#### Sequence Listing

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<211> 249
<212> DNA
<213> Homo sapiens
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ctgatggtgg aggaagggc gtctatcaat caatcactgt tgctgttatc 150
acatgcaagt atccagaggc tottgagcaa ggcagagggg atcccattta 200
tttgggaatc cagaatccag aaatgtgttt gtattgtgag aaggttgga 249
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<211> 468
<212> DNA
<213> Homo sapiens
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ggtgctttat ctgcataata accagcttct agctggaggg ctgcatgcag 100
ggaaggtcat taaaggtgaa gagatcagcg tggtccccaa tcggtggctg 150
gatgccagcc tgtcccccgt catcctgggt gtccagggtg gaagccagtg 200
cctqtcatqt qqqqtqqqqc aqqaqccqac tctaacacta qaqccaqtqa 250
acatcatgga gctctatctt ggtgccaagg aatccaagag cttcaccttc 300
tacoggoggg acatggggct cacctccage ttegagtegg etgectacce 350
gggetggttc ctgtgcacgg tgcctgaagc cgatcagcct gtcagactca 400
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ttccagcagt gtgactag 468
<210> 13
<211> 155
<212> PRT
<213> Homo sapiens
<400> 13
Met Val Leu Ser Gly Ala Leu Cys Phe Arg Met Lys Asp Ser Ala
Leu Lys Val Leu Tyr Leu His Asn Asn Gln Leu Leu Ala Gly Gly
Leu His Ala Gly Lys Val Ile Lys Gly Glu Glu Ile Ser Val Val
Pro Asn Arg Trp Leu Asp Ala Ser Leu Ser Pro Val Ile Leu Gly
Val Gln Glv Glv Ser Gln Cvs Leu Ser Cvs Glv Val Glv Gln Glu
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65

<400> 14

<223> unknown base

00 g,

m On

0

getecegeea ggagaaagga acattetgag gggagtetae accetgtgga 50 getcaagatg gteetgagtg gggegetgtg etteegaatg aaggactegg 100 cattgaaggt getttatetg cataataace agettetage tggagggetg 150 catgcaggga aggtcattaa aggtgaagag atcagcgtgg tccccaatcg 200 gtggctggat gccagcctgt cccccgtcat cctgggtgtc cagggtggaa 250 gecagtgeet gtcatgtggg gtggggcagg agnegaetet aacat 295

<210> 15 <211> 1385

<212> DNA

<213> Mus musculus

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agtotgtocc ctgtcatcct gggcgttcaa ggaggaagcc agtgcctatc 350 ttqtqqqaca qaqaaaqqqc caattctqaa acttqaqcca qtqaacatca 400 tggageteta ceteggggee aaggaateaa agagetteae ettetaeegg 450 egggatatgg gtettacete cagettegaa teegetgeet acceaggetg 500 gttcetctgc acctcaccgg aagetgacca gcctgtcagg ctcactcaga 550 teectgagga cecegeetgg gatgeteeca teacagaett etaettteag 600 cagtgtgact agggctgcgt ggtccccaaa actccataag cagaggcaga 650 gtaggcagtg geggeteetg atagaggata gagagacaga ggageteeac 700 agtaggtggc ttactcctct cettcectac tggactcccg ettctgacet 750 aaggcacaca gacactetet teteetgcat eccagtgetg gtaaatette 800 tggtatttgg agctcaatgt gtagattett teagattgga tggtaetace 850 tctggtgtgg aacccaatag aaaccacgta ggaccaacaa agagcaacat 900 aaaagattot tgggtgaaga agaggtggga actgttcata catagtaaga 950 totgacacag tacotcagaa gtootgocat toottatgtt otggagaaag 1000 tggaggggg gtcaccaaga ctttctctgg ctggctgggc cctttccctc 1050 aacettetg acatetgeag ceteteteat tettgeette attetetgge 1100 cctgaaccga gagggtgata tcaggatagc tgacagaaga tgaccaggca 1150 cactgtcctg gtttgaaacc agaggggaca ataaaaaacc ctgattctgg 1200 tototactca cataaaaaga agottgtgaa cattaagtgg gaagagattg 1250 ctactaaata acatacettg taatttcate ttaattaaaa tatacttete 1300 aaaaacatgc ggccgcaagc ttattccatt tagga 1385

<210> 16 <211> 155

<212> PRT

<213> Mus musculus

<400> 16

Met Val Leu Ser Gly Ala Leu Cys Phe Arg Met Lys Asp Ser Ala 1 5 10 15

Leu Lys Val Leu Tyr Leu His Asn Asn Gln Leu Leu Ala Gly Gly  $20 \\ 25 \\ 30$ 

Leu His Ala Glu Lys Val Ile Lys Gly Glu Glu Ile Ser Val Val

Pro Asn Arg Ala Leu Asp Ala Ser Leu Ser Pro Val Ile Leu Gly  $50 \hspace{1cm} 55 \hspace{1cm} 60 \hspace{1cm}$ 

```
Val Gln Gly Gly Ser Gln Cys Leu Ser Cys Gly Thr Glu Lys Gly For Ile Leu Lys Leu Glu Pro Val Asn Ile Met Glu Leu Tyr Leu
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Gly Ala Lys Glu Ser Lys Ser Phe Thr Phe Tyr Arg Arg Asp Met

Gly Leu Thr Ser Ser Phe Glu Ser Ala Ala Tyr Pro Gly Trp Phe

Leu Cys Thr Ser Pro Glu Ala Asp Gln Pro Val Arg Leu Thr Gln

Phe Gln Gln Cys Asp

<210> 17

<211> 382

<212> DNA

<213> Mus musculus

<400> 17

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atgatggete tgagtgggge actatgette egaatgaagg atteageett 150
gaaggtactg tatetgeaca ataaceaget getggetgga ggaetgeacg 200
cagagaaggt cattaaaggt gaggagatea gtgttgteee aaategggea 250
ctggatgeea geetgteeee tgteateetg ggegtteaag gaggaageea 300
gtgeetatet tgtgggacag agaaagggee aattetgaaa ettgageeag 350
tgaacateat ggagetetae eteggggeea ag 382

<210> 18

<211> 626 <212> DNA

<213> Homo sapiens

<400> 18

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attitigtica cacaagteca aaggtgaaga actiaaacce gaagaaatte 150
agcatteatg accaggatea caaagtactg gteetggact etgggaatet 200
catageagtt ecagataaaa actacataeg eccagagate ttettigeat 250
tageeteate ettgagetea geetetgegg agaaaggaag teegattete 300

ctgggggtct ctaaagggga gttttgtctc tactgtgaca aggataaagg 350
acaaagtcat ccatccettc agctgaagaa ggagaaactg atgaagctgg 400
ctgcccaaaa ggaatcagca cgccggccct tcatctttta tagggctcag 450
gtgggctcct ggaacatgct ggagtcggcg gctcaccccg gatggtcat 500
ctgcacctcc tgcaattgta atgagcctgt tggggtgaca gataaatttg 550
agaacaggaa acacattgaa ttttcattc aaccagtttg caaagctgaa 600
atgagccca gtgaggtcag cgatta 626

<210> 19

<211> 207

<212> PRT

<213> Homo sapiens

<400> 19

Met Gly Ser Glu Asp Trp Glu Lys Asp Glu Pro Gln Cys Cys Leu 1  $\phantom{\bigg|}$  10  $\phantom{\bigg|}$  15

Ala Met Asn Phe Val His Thr Ser Pro Lys Val Lys Asn Leu Asn \$35\$ 40 45

Pro Lys Lys Phe Ser Ile His Asp Gln Asp His Lys Val Leu Val 50 60

Leu Asp Ser Gly Asn Leu Ile Ala Val Pro Asp Lys Asn Tyr Ile
65 70 75

Arg Pro Glu Ile Phe Phe Ala Leu Ala Ser Ser Leu Ser Ser Ala 80 85 90

Ser Ala Glu Lys Gly Ser Pro Ile Leu Leu Gly Val Ser Lys Gly 95 \$100\$

Lys Glu Ser Ala Arg Arg Pro Phe Ile Phe Tyr Arg Ala Gln Val 140 145 150

Gly Ser Trp Asn Met Leu Glu Ser Ala Ala His Pro Gly Trp Phe 155 160 165

Lys Phe Glu Asn Arg Lys His Ile Glu Phe Ser Phe Gln Pro Val 185 190 195

Cys Lys Ala Glu Met Ser Pro Ser Glu Val Ser Asp

<210> 20 <211> 506 <212> DNA

<213> Homo sapiens

<400> 20

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attitigtica cacaaagate tietitigeat tageeteate etigagetea 150
geetetgegg agaaaggaag teegatiete etgaggete etaaagggga 200
gittigtete taetgtgaea aggataaagg acaaagteat ceatecette 250
agetgaagaa ggagaaaetg atgaagetgg etgeecaaa ggaateagea 300
egeeggeet teatetitia tagggeteag gigggeteet ggaacatget 350
ggagteggeg geteaceeeg gatggteat etgeacetee tgeaatigta 400
atgageetgt tggggtgaea gataaattig agaacaggaa acacattgaa 450
titteatite aaceagtitg caaagetgaa atgageeeca gtgaggteag 500
egatta 506

<210> 21 <211> 167 <212> PRT

<213> Homo sapiens

<400> 21

Met Gly Ser Glu Asp Trp Glu Lys Asp Glu Pro Gln Cys Cys Leu 1 5 10 15

Glu Asp Pro Ala Val Ser Pro Leu Glu Pro Gly Pro Ser Leu Pro

Ala Met Asn Phe Val His Thr Lys Ile Phe Phe Ala Leu Ala Ser  $35 \hspace{1cm} 40 \hspace{1cm} 45$ 

Ser Leu Ser Ser Ala Ser Ala Glu Lys Gly Ser Pro Ile Leu Leu  $50 \\ 55 \\ 60$ 

Gly Val Ser Lys Gly Glu Phe Cys Leu Tyr Cys Asp Lys Asp Lys

Lys Leu Ala Ala Gln Lys Glu Ser Ala Arg Arg Pro Phe Ile Phe 95  $$100\,$ 

Tyr Arg Ala Gln Val Gly Ser Trp Asn Met Leu Glu Ser Ala Ala 110 \$115\$

His Pro Gly Trp Phe Ile Cys Thr Ser Cys Asn Cys Asn Glu Pro

Val Gly Val Thr Asp Lys Phe Glu Asn Arg Lys His Ile Glu Phe 140 145 150

Ser Asp

<210> 22

<211> 561 <212> DNA

<213> Homo sapiens

<400> 22

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agatcttott tgcattagce toatcottga gctcagcott tgcggagaaa 150
ggaagtccga ttotcctggg ggtotctaaa ggggagttt gtotctactg 200
tgacaaggat aaaggacaaa gtcatccatc cottcagctg aagaaggaga 250
aactgatgaa gctggotgoc caaaaggaat cagcagccg gccottcatc 300
ttttataggg otcaggtggg otcctgaaa tggtaggat cggcggotca 350
ccccggatgg ttoatctgaa cotcctgaa ttgtaatgag cctgttgggg 400
tgacagataa atttgagaac aggaaacaca ttgaatttc atttcaacca 450
gtttgcaaag otgaaatgag occcagtgag gtcagcgatt aggaaactgc 500
cccattgaac gccttcctcq ctaatttgaa caaatggta aaaaacacca 550

<210> 23 <211> 561

<212> DNA

<213> Homo sapiens

aacctgetca c 561

<400> 23

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cetgagacce ttagagtate gteaaggtet atttttgatg tatgegggte 100
tetagaagaa acgtaategg agtaggaact egagteggag acgeetettt 150
cetteagget aagaggacce eeagagattt eeeetaaaa eagagatgae 200
actgtteeta ttteetgttt eagtaggtag ggaagtegae ttetteetet 250
ttgaetaett egaeegaegg gttteetta gteggegge egggaagtag 300
aaaatateee gagteeaeee gaggaeettg taegaeetea geegeegagt 350

ggggcctacc aagtagacgt ggaggacgtt aacattactc ggacaacccc 400 actgtctatt taaactcttg tootttgtgt aacttaaaag taaagttggt 450 caaacgtttc gactttactc ggggtcactc cagtcgctaa tcctttgacg 500 gggtaacttg cggaaggagc gattaaactt gattaacata tttttgtggt 550 ttggacgagt g 561

<210> 24 <211> 839

<212> DNA

<213> Homo sapiens

<400> 24 ggccctcgag gccaagaatt cggcacgagg cttcattcca ttttctgttg 50

agtaataaac tcaacgttga aaatgtcctt tgtgggggag aactcaggag 100 tgaaaatggg etetgaggae tgggaaaaag atgaacccca gtgctgetta 150 gaagacccqq ctqqaaqccc cctqqaacca gqcccaagcc tccccaccat 200 gaattttgtt cacacaagtc caaaggtgaa gaacttaaac ccgaagaaat 250 tcagcattca tgaccaggat cacaaagtac tggtcctgga ctctgggaat 300 ctcatagcag ttccagataa aaactacata cgcccagaga tcttctttgc 350 attagectea teettgaget cageetetge ggagaaagga agteegatte 400 tectgggggt etetaaaggg gagttttgte tetactgtga caaggataaa 450 ggacaaagte atecateeet teagetgaag aaggagaaac tgatgaaget 500 ggctgcccaa aaggaatcag cacgccggcc cttcatcttt tatagggctc 550 aggtgggete etggaacatg etggagtegg eggeteacce eggatggtte 600 atctgcacct cctgcaattg taatgagcct gttggggtga cagataaatt 650 tgagaacagg aaacacattg aattttcatt tcaaccagtt tgcaaagctg 700 aaatqaqccc caqtqaqqtc agcqattaqq aaactqcccc attqaacqcc 750 ttectegeta atttgaacta attgtataaa aacaccaaac etgeteacta 800

aaaaaaaaa aaaaaaacgt ttgcggccgc aagcttatt 839

<sup>&</sup>lt;210> 25

<sup>&</sup>lt;211> 218

<sup>&</sup>lt;212> PRT

<sup>&</sup>lt;213> Homo sapiens

<sup>&</sup>lt;400> 25

Met Ser Phe Val Gly Glu Asn Ser Gly Val Lys Met Gly Ser Glu

Asp Trp Glu Lys Asp Glu Pro Gln Cys Cys Leu Glu Asp Pro Ala

```
Gly Ser Pro Leu Glu Pro Gly Pro Ser Leu Pro Thr Met Asn Phe
Val His Thr Ser Pro Lys Val Lys Asn Leu Asn Pro Lys Lys Phe
Ser Ile His Asp Gln Asp His Lys Val Leu Val Leu Asp Ser Gly
Asn Leu Ile Ala Val Pro Asp Lys Asn Tyr Ile Arg Pro Glu Ile
Phe Phe Ala Leu Ala Ser Ser Leu Ser Ser Ala Ser Ala Glu Lys
Gly Ser Pro Ile Leu Leu Gly Val Ser Lys Gly Glu Phe Cys Leu
Tyr Cys Asp Lys Asp Lys Gly Gln Ser His Pro Ser Leu Gln Leu
Lys Lys Glu Lys Leu Met Lys Leu Ala Ala Gln Lys Glu Ser Ala
                                    145
Arg Arg Pro Phe Ile Phe Tyr Arg Ala Gln Val Gly Ser Trp Asn
Met Leu Glu Ser Ala Ala His Pro Gly Trp Phe Ile Cys Thr Ser
Cys Asn Cys Asn Glu Pro Val Gly Val Thr Asp Lys Phe Glu Asn
Arg Lys His Ile Glu Phe Ser Phe Gln Pro Val Cys Lys Ala Glu
                                    205
Met Ser Pro Ser Glu Val Ser Asp
<210> 26
<211> 177
<212> PRT
<213> Homo sapiens
<400> 26
Met Glu Ile Cys Arg Gly Leu Arg Ser His Leu Ile Thr Leu Leu
Leu Phe Leu Phe His Ser Glu Thr Ile Cys Arg Pro Ser Gly Arg
Lys Ser Ser Lys Met Gln Ala Phe Arg Ile Trp Asp Val Asn Gln
Lys Thr Phe Tyr Leu Arg Asn Asn Gln Leu Val Ala Gly Tyr Leu
Gln Gly Pro Asn Val Asn Leu Glu Glu Lys Ile Asp Val Val Pro
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```
Ile Glu Pro His Ala Leu Phe Leu Gly Ile His Gly Gly Lys Met
 Cys Leu Ser Cys Val Lys Ser Gly Asp Glu Thr Arg Leu Gln Leu
Glu Ala Val Asn Ile Thr Asp Leu Ser Glu Asn Arg Lys Gln Asp
 Lys Arg Phe Ala Phe Ile Arg Ser Asp Ser Gly Pro Thr Thr Ser
 Phe Glu Ser Ala Ala Cys Pro Gly Trp Phe Leu Cys Thr Ala Met
Glu Ala Asp Gln Pro Val Ser Leu Thr Asn Met Pro Asp Glu Gly
Val Met Val Thr Lys Phe Tyr Phe Gln Glu Asp Glu
<210> 27
<211> 169
<212> PRT
<213> Homo sapiens
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Tyr Gln Ser Met Cys Lys Pro Ile Thr Gly Thr Ile Asn Asp Leu
Asn Gln Gln Val Trp Thr Leu Gln Gly Gln Asn Leu Val Ala Val
Pro Arg Ser Asp Ser Val Thr Pro Val Thr Val Ala Val Ile Thr
Cys Lys Tyr Pro Glu Ala Leu Glu Gln Gly Arg Gly Asp Pro Ile
Tyr Leu Gly Ile Gln Asn Pro Glu Met Cys Leu Tyr Cys Glu Lys
Val Gly Glu Gln Pro Thr Leu Gln Leu Lys Glu Gln Lys Ile Met
Asp Leu Tyr Gly Gln Pro Glu Pro Val Lys Pro Phe Leu Phe Tyr
Arg Ala Lys Thr Gly Arg Thr Ser Thr Leu Glu Ser Val Ala Phe
Pro Asp Trp Phe Ile Ala Ser Ser Lys Arg Asp Gln Pro Ile Ile
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Asn Ile Asn Asp

Leu Thr Ser Glu Leu Gly Lys Ser Tyr Asn Thr Ala Phe Glu Leu

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<210> 28
<211> 167
<212> PRT
<213> Homo sapiens
<400> 28
Met Gly Ser Glu Asp Trp Glu Lys Asp Glu Pro Gln Cys Cys Leu
 Glu Asp Pro Ala Gly Ser Pro Leu Glu Pro Gly Pro Ser Leu Pro
 Thr Met Asn Phe Val His Thr Lys Ile Phe Phe Ala Leu Ala Ser
 Ser Leu Ser Ser Ala Ser Ala Glu Lys Gly Ser Pro Ile Leu Leu
 Gly Val Ser Lys Gly Glu Phe Cys Leu Tyr Cys Asp Lys Asp Lys
 Gly Gln Ser His Pro Ser Leu Gln Leu Lys Lys Glu Lys Leu Met
 Lys Leu Ala Ala Gln Lys Glu Ser Ala Arg Arg Pro Phe Ile Phe
 Tyr Arg Ala Gln Val Gly Ser Trp Asn Met Leu Glu Ser Ala Ala
 His Pro Gly Trp Phe Ile Cys Thr Ser Cys Asn Cys Asn Glu Pro
 Val Gly Val Thr Asp Lys Phe Glu Asn Arg Lys His Ile Glu Phe
 Ser Phe Gln Pro Val Cys Lys Ala Glu Met Ser Pro Ser Glu Val
                                     160
 Ser Asp
<210> 29
<211> 31
<212> DNA
<213> Homo sapiens
<400> 29
ggeggateca aaatgggete tgaggaetgg g 31
<210> 30
<211> 30
<212> DNA
<213> Homo sapiens
<400> 30
geggaattet aategetgae eteaetgggg 30
<210> 31
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<211> 9
<212> PRT
<213> Homo sapiens
<4400> 31
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    1
    5
<210> 32
<211> 10
<212> PRT
<213> Homo sapiens
<4400> 32
    Val Leu Ser Gly Ala Leu Cys Phe Arg Met
    1
    5
    10
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